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PATENT APPLICATION TRANSMITTAL LETTER

Docket No.:
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98126-CIP

To the Assistant Commissioner for Patents:

Transmitted herewith for filing under 35 U.S.C. 111 and 37 C.F.R. 1.53 is the Continuation-in-part patent application of: Brad A. Ozenberger, J.S. Jacobsen, Eileen Kajkowski, J. Bard and S. Walker

entitled: "β-AMYLOID PEPTIDE-BINDING PROTEINS AND
POLYNUCLEOTIDES ENCODING THE SAME"

Enclosed are:

10 informal sheet(s) of drawings.
 an assignment of the invention to _____
 A certified copy of a _____ application.
 Declaration Signed Unsigned
 Power of Attorney
 Information Disclosure Statement
 Preliminary Amendment
 Other:

CLAIMS AS FILED

	Total No. Filed:	No. Allowed:	No. Extra:	Rate	Fee
Total Claims	24	-20 =	4	x \$22.00	\$ 88.00
Independent claims	14	- 3 =	11	x \$82.00	\$902.00
Multiple Dependent Claims Present				x \$270.00	
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Dated: October 14, 1998

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**β-AMYLOID PEPTIDE-BINDING PROTEINS
AND POLYNUCLEOTIDES ENCODING THE SAME**

5 This application claims benefit of U.S. Ser. No. 09/060, 609 filed April 15, 1998 and U.S. Provisional Application 60/064,583, filed April 16, 1997, the content of which is incorporated into this application by reference.

10 **Field of the Invention**

 The present invention relates to a novel polynucleotides and proteins encoded by such polynucleotides, along with therapeutic, diagnostic, and research utilities for these polynucleotides and proteins. In particular, the invention relates to polynucleotides and proteins encoded 15 by such polynucleotides which bind to β-amyloid peptide, one of the primary components of amyloid deposits associated with Alzheimer's Disease.

Background of the Invention

 Alzheimer's disease (AD) is a progressive dementing disorder of 20 the elderly characterized by a series of structural abnormalities of the brain. Neurons in multiple regions of the central nervous system (CNS) become dysfunctional and die, resulting in alterations in synaptic inputs. Cell bodies and proximal dendrites of these vulnerable neurons contain 25 neurofibrillary tangles composed of paired helical filaments, the main component of which is a phosphorylated microtubular-binding protein, namely tau. One of the hallmarks of the disease is the accumulation of amyloid containing deposits within the brain called senile (or neuritic) plaques. The principal component of amyloid plaques is β-amyloid peptide (hereinafter "BAP", also referred in the literature as Aβ, βAP, etc.) which 30 forms dense aggregates during the course of AD.

 BAP is a 39-43 amino acid peptide derived by proteolytic cleavage of amyloid precursor protein (hereinafter "APP") and composed of a

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portion of the transmembrane domain and the luminal/extracellular domain of APP. It is thought that the BAP peptide comprising 42 amino acids (BAP42) is potentially the more toxic aggregated form in humans. APP occurs as several BAP-containing isoforms. The major forms are

5 comprised of 695, 751, and 770 amino acids, with the latter two APP containing a domain that shares structural and functional homologies with Kunitz serine protease inhibitors. In normal individuals, BAP does not accumulate and is rapidly removed from circulating fluids. However, the peptide can form plaques on surfaces of dystrophic dendrites and axons, 10 microglia, and reactive astrocytes. The aggregation and deposition of BAP in neuritic plaques is postulated as one of the initiating events of AD. Investigation of the events leading to the expression and consequences of BAP and their individual roles in AD is a major focus of neuroscience research. In particular, the discovery of proteins that bind BAP is critical 15 to advance understanding of the pathogenesis of the disease and to potentially introduce novel therapeutic targets.

Until the present invention, proteins and fragments thereof which bind with human BAP and which may be involved in the biological effects of BAP in AD had not been identified.

20 **Summary of the Invention**

This invention provides novel isolated polynucleotides which encode gene products that selectively bind human β -amyloid peptide (BAP) amino acid sequences.

In one embodiment, the present invention provides a composition 25 comprising an isolated polynucleotide selected from the group consisting of:

(a) polynucleotide comprising the nucleotide sequence of SEQ ID NO: 1;

(b) a polynucleotide comprising the nucleotide sequence of a β -30 amyloid peptide-binding protein (BBP) of clone BBP1-fl deposited under accession number ATCC 98617;

(c) a polynucleotide encoding a β -amyloid peptide-binding protein (BBP) encoded by the cDNA insert of clone BBP1-fl deposited under accession number ATCC 98617;

(d) a polynucleotide comprising the nucleotide sequence of SEQ ID NO: 1 from nucleotide 202 to nucleotide 807;

(e) a polynucleotide comprising the nucleotide sequence of a β -amyloid peptide-binding protein (BBP) of clone pEK196 deposited under accession number ATCC 98399;

(f) a polynucleotide encoding a β -amyloid peptide-binding protein (BBP) encoded by the cDNA insert of clone pEK196 deposited under accession number ATCC 98399;

(g) a polynucleotide encoding a protein comprising the amino acid sequence of SEQ ID NO: 2;

(h) a polynucleotide encoding a protein comprising a fragment of the amino acid sequence of SEQ ID NO: 2 having human β -amyloid peptide binding activity, the fragment comprising the amino acid sequence from amino acid 68 to amino acid 269 of SEQ ID NO: 2;

(i) a polynucleotide which is an allelic variant of the polynucleotide of (a)-(f) above;

(j) a polynucleotide which encodes a species homologue of the protein of (g)-(i) above; and

(l) a polynucleotide capable of hybridizing under stringent conditions to any one of the polynucleotides specified in (a)-(h).

Preferably such polynucleotide comprises the nucleotide sequence of SEQ ID NO: 1; the nucleotide sequence of a β -amyloid peptide-binding protein (BBP) of clone BBP1-fl deposited under accession number ATCC 98617; or a polynucleotide encoding a β -amyloid peptide-binding protein (BBP) encoded by the cDNA insert of clone BBP1-fl deposited under accession number ATCC 98617. Another embodiment provides the gene corresponding to the cDNA sequence of SEQ ID NO: 1.

In other embodiments, the present invention provides a composition comprising a protein, wherein said protein comprises an amino acid sequence selected from the group consisting of:

- (a) the amino acid sequence of SEQ ID NO: 2;
- 5 (b) the amino acid sequence of SEQ ID NO: 2 from amino acid 68 to amino acid 269;
- (c) the amino acid sequence encoded by the cDNA insert of clone BBP1-fl deposited under accession number ATCC 98617; and
- (d) 10 fragments of the amino acid sequence of SEQ ID NO: 2 comprising the amino acid sequence from amino acid 185 to amino acid 217 of SEQ ID NO: 2.

Preferably such protein comprises the amino acid sequence of SEQ ID NO: 2 or the amino acid sequence of SEQ ID NO: 2 from amino acid 68 to amino acid 269. Fusion proteins are also claimed in the present 15 invention.

In certain preferred embodiments, the polynucleotide is operably linked to an expression control sequence. The invention also provides a host cell, including bacterial, yeast, insect, and mammalian cells, transformed with such polynucleotides compositions.

20 Processes are also provided for producing a BBP which comprises

- (a) growing a culture of the host cell of claim 3 in a suitable culture medium; and (b) purifying the protein from the culture medium.

Compositions comprising an antibody which specifically reacts with such BBPs are also provided by the present invention.

25 Methods and diagnostic processes are provided for detecting a disease state characterized by the aberrant expression of human BAP, as well as methods for identifying compounds which regulate the activity of BBPs.

Another embodiment of the invention includes transgenic animals 30 comprising a polynucleotide encoding a BBP operably linked to an expression control sequence.

Brief Description of the Drawings

The following drawings depict certain embodiments of the invention. They are illustrative only and do not limit the invention otherwise disclosed herein.

5 Figure 1: Yeast 2-hybrid screen design. A Y2H host strain expressing the Gal4 DNA-binding domain fused to BAP₄₂ (BAP^{BD}; plasmid containing TRP1 marker) and nonfusion BAP₄₂ (BAP; plasmid containing URA3 marker) was transformed with a Y2H human fetal brain cDNA library (plasmid containing LEU2 marker) expressing Gal4 activation

10 domain fusion proteins (unknown^{AD}) as described. Therefore, strains contained three episomal plasmids, denoted by circles, expressing the indicated protein. Positive protein-protein interactions reconstituted Gal4 activity at the upstream activating sequence (GALUAS) thereby inducing transcription of the reporter gene HIS3.

15 Figure 2: Demonstration of BBP1/BAP association. Y2H strains were assayed for histidine prototrophy by making 10-fold serial dilutions and spotting 5 μ l on synthetic agar medium lacking tryptophan, leucine, histidine and containing 25 mM 3-amino-triazole as described. All strains contain the BAP fusion protein expression plasmid pEK162 as indicated

20 by the label BAP. The first columns (vector) contain independently derived strains carrying pEK162 and the vector pACT2 expressing an irrelevant fusion protein. These serve as a measure of background for comparison with strains expressing target proteins. The columns marked by BBP1Dtm express a truncated BBP1 from pEK198, as described in the

25 text. The interaction between BAP and BBP1 Δ tm fusion proteins reconstitutes Gal4 activity, resulting in induction of a HIS3 reporter gene (see Figure 1), observed as enhanced prototrophic growth compared to the control strains.

30 Figure 3: Bioassays demonstrating BBP1 interactions with G α proteins. The predicted intracellular domain of BBP1 was expressed as a Gal4 DNA-binding domain with portions of rat G α s, G α o, or G α i2

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expressed as Gal4 activation domain fusion proteins. Y2H responses of two independently derived clones of each strain were compared to responses of cells lacking a G protein component (vector). The protocol is as described in the legend to Figure 2.

5 Figure 4: Localization of the interactions between BBP1 and BAP.

BBP1 Δ tm was divided into two overlapping segments as described in the text. These proteins, BBP1 Δ C or BBP1 Δ N, were assayed for interactions with BAP. The assay method and the strains labeled vector or BBP1 Δ tm are as described in the legend to Figure 2. Strains labeled BBP1 Δ C or

10 BBP1 Δ N express the indicated BBP1 segment as a fusion protein.

Figure 5: Expression of BBP1 mRNA in human tissues (A) and brain regions (B). Nylon membranes blotted with 2 μ g size fractionated poly-A RNA isolated from the indicated tissues were obtained from CLONTECH. These were hybridized with a radiolabeled BBP1 cDNA probe 15 as described. A predominant band corresponding to 1.25 kb (determined from molecular weight markers, not shown) was observed in all lanes. Higher molecular weight bands likely correspond to heteronuclear RNA; the BBP1 gene contains several introns. Blots were stripped and reprobed with β -actin as a loading and RNA integrity control; all lanes exhibited 20 equivalent signal (data not shown).

Figure 6: Expression of BBP1 and APP in cells of the hippocampus. Images of in situ hybridization autoradiograms showing the pattern of BBP1 (A) and APP (B) expression in human hippocampal and entorhinal cortex. The sections used to generate these images were 25 taken from postmortem specimens obtained from two different patients. Abbreviations: DG = dentate gyrus; CA1 = hippocampal subfield; EC = entorhinal cortex.

Figure 7: Comparison of BBP1 interactions with human or rodent BAP. Rodent BAP was engineered and expressed as a fusion protein as 30 described in the text. The strains labeled human BAP are identical to those shown in Figure 2. The strains labeled rodent BAP express rodent

BBP1 Δ tm = BBP1 Δ tm

BAP as the Gal4 DNA-binding domain fusion. Vector indicates control strains containing only vector opposing the BAP fusion proteins; BBP1 indicates strains expressing the BBP1 Δ tm fusion protein.

Fig. 8. In vitro BBP1 binding to beta-amyloid protein (1-42). In vitro transcribed and translated N-terminally myc-tagged BBP1 (~20ng) was incubated with disaggregated human [125 I]-Tyr-BAP₍₁₋₄₂₎. The myc-tagged BBP1 was immunoprecipitated with mouse anti-myc antibody and rabbit anti-mouse IgG antibody conjugated to protein A-Agarose beads. lane 1, MW markers; lane2, 4, 6, 8: blank; lane 3: [125 I]-Tyr-BAP₍₁₋₄₂₎(disaggregated; \approx 25nCi); lane 5: Pre-incubation of rabbit anti-mouse IgG antibody conjugated to protein A-Agarose beads in Superblock, plus mouse anti-myc antibody and radiolabeled BAP (no BBP1); lane 7: Pre-incubation of rabbit anti-mouse IgG antibody conjugated to protein A-Agarose beads in low salt binding buffer (see Material and Methods), plus mouse anti-myc antibody and radiolabeled BAP (no BBP1); lane9: Pre-incubation of rabbit anti-mouse IgG antibody conjugated to protein A-Agarose beads in Superblock, plus mouse anti-myc antibody, radiolabeled BAP, and in vitro synthesized myc-tagged BBP1.

Fig. 9. Effect of BBP1 on Nt2 Stem Cells Exposed to BAP. Nt2 neurons, and Nt2 stem cells transfected with pEGFP alone or cotransfected with pEGFP plus pBBP1 (also referred to as pOZ363), were treated for 48 hrs with aged (toxic) or fresh (nontoxic) BAP preparations as described in the text. Final concentrations were 2 μ M or 5 μ M, respectively. Cells were prepared for determinations of nuclear morphology as described. Values represent the mean percent condensed nuclei, derived from viewing multiple fields for each sample. For the stem cell populations, only transfected cells (EGFP +) were scored.

Fig. 10. Sensitivity of Nt2 stem cells to modified BBP. Nt2 stem cells were transfected with pEGFP alone or cotransfected with pEGFP plus BBP1 or BBP1(R138E) expression plasmids. Samples were treated with BAP at the indicated concentrations for 44 hrs. Cells were prepared

for determinations of nuclear morphology as described. Values represent percent condensed nuclei of transfected cells (EGFP +). The differences between values for BBP1 samples versus controls at 2 or 8 μ M BAP are highly significant ($P < 0.005$) as determined by a Yates modified chi-

5 square test of probability.

Detailed Description of the invention

The present invention relates to the isolation and cloning of a human β -amyloid peptide binding protein (BBP1). BBP1 has been characterized as a fusion protein in a yeast 2 hybrid assay as binding to

10 BAP, specifically the 42 amino acid fragment of BAP (BAP42).

Expression of BBP1 has been shown in human tissues and in specific brain regions (Figure 5). Importantly BBP1 has been demonstrated to selectively bind human BAP in a yeast 2 hybrid system as compared to rodent BAP. These findings support the premise that the BBP1 of the

15 present invention may be used in the diagnosis and treatment of Alzheimer's Disease.

The BBP1 Coding Sequence

The initial human BBP1 clone (designated clone 14) was obtained 20 by using a yeast 2-hybrid (Y2H) genetic screen developed to identify proteins which interact with human BAP₄₂, a potentially more toxic form of BAP. BAP₄₂ was expressed fused to the yeast Gal4 DNA-binding domain and was also expressed as free peptide (Figure 1). This strain was transformed with a human fetal brain cDNA Y2H library. A single

25 clone, denoted #14, from approximately 10⁶ independent transformants, produced consistent reporter gene activation and contained a substantial open reading frame continuous with that of the GAL4 domain. The cDNA insert comprised 984 base pairs, terminating in a poly-A tract. This sequence encoded 201 amino acids (amino acid 68 to amino acid 269 of

30 SEQ ID NO: 2) with two regions of sufficient length and hydrophobicity to transverse a cellular membrane. There are also potential asparagine-linked

65 66 67 68 69 70 71 72 73 74 75 76 77 78 79 80

glycosylation sites. Clone 14 was designated clone pEK196 and deposited as ATCC 98399 .

The library-derived plasmid was isolated from clone 14 and used to reconstruct Y2H assay strains. Examination of these strains

- 5 demonstrated that the BAP fusion protein specifically interacted with the clone 14 protein, although the response was weak. Since protein domains of strong hydrophobicity, such as transmembrane regions, inhibit Y2H responses (Ozenberger, unpublished data), clone 14 insert was truncated (BBP1 Δ tm; see Table 2 below for further description) to remove
- 10 the region of strongest hydrophobicity and retested for interactions with BAP. A much more robust Y2H response was observed with BBP1 Δ tm, supporting the notion that the deleted sequences encode a potential transmembrane ("tm") anchor. Clone 14 identifies a novel BAP binding protein in the form of a fusion protein.
- 15 The BBP1 cDNA sequences contained in clone 14 were identified as lacking the 5' end of the protein coding region as no potential initiating methionine codon was present. Multiple attempts at conventional 5' RACE (rapid amplification of cDNA ends) utilizing a standard reverse-transcriptase only resulted in the addition of 27 nucleotides. Thus, a
- 20 genomic cloning approach as described in Example 2, below, was used to isolate the 5' terminus.

Since the 5' coding sequence terminus was derived from a genomic library, there existed the possibility that this region contained introns. This potentiality was investigated by two methods as described

- 25 in Example 2, below. The resulting data confirmed the upstream sequences (both from genomic and cDNA sources) and the lack of introns in this region. Plasmid BBP1-f1 containing a cDNA insert encoding the full length BBP1 protein coding region was deposited in the American Type Culture Collection with accession number 98617. The entire coding
- 30 region and deduced protein sequence is shown in SEQ ID NOS:1 and

SEQUENCE LISTING

2..The 3' nontranslated nucleotide sequences are contained in the original clone 14 (pEK196).

In accordance with the present invention, nucleotide sequences which encode BBP1, fragments, fusion proteins or functional equivalents

5 thereof, may be used to generate recombinant DNA molecules that direct the expression of BBP1, or a functionally active peptide, in appropriate host cells. Alternatively, nucleotide sequences which hybridize to portions of the BBP1 sequence may be used in nucleic acid hybridization assays, Southern and Northern blot assays, etc.

10 The invention also includes polynucleotides with sequences complementary to those of the polynucleotides disclosed herein.

The present invention also includes polynucleotides capable of hybridizing under reduced stringency conditions, more preferably stringent conditions, and most preferably highly stringent conditions, to

15 polynucleotides described herein. Examples of stringency conditions are shown in the table below: highly stringent conditions are those that are at least as stringent as, for example, conditions A-F; stringent conditions are at least as stringent as, for example, conditions G-L; and reduced stringency conditions are at least as stringent as, for example, conditions

20 M-R.

Stringency Conditions

Stringency Condition	Polynucleotide Hybrid	Hybrid Length (bp) ¹	Hybridization Temperature and BufferH	Wash Temperature and BufferH
A	DNA:DNA	> 50	65EC; 1xSSC -or- 42EC; 1xSSC, 50% formamide	65EC; 0.3xSSC
B	DNA:DNA	< 50	T _B *; 1xSSC	T _B *; 1xSSC
C	DNA:RNA	> 50	67EC; 1xSSC -or- 45EC; 1xSSC, 50% formamide	67EC; 0.3xSSC
D	DNA:RNA	< 50	T _D *; 1xSSC	T _D *; 1xSSC

E	RNA:RNA	\$ 50	70EC; 1xSSC -or- 50EC; 1xSSC, 50% formamide	70EC; 0.3xSSC
F	RNA:RNA	<50	T _F *; 1xSSC	T _f *; 1xSSC
G	DNA:DNA	> 50	65EC; 4xSSC -or- 42EC; 4xSSC, 50% formamide	65EC; 1xSSC
H	DNA:DNA	<50	T _H *; 4xSSC	T _h *; 4xSSC
I	DNA:RNA	> 50	67EC; 4xSSC -or- 45EC; 4xSSC, 50% formamide	67EC; 1xSSC
J	DNA:RNA	<50	T _J *; 4xSSC	T _j *; 4xSSC
K	RNA:RNA	> 50	70EC; 4xSSC -or- 50EC; 4xSSC, 50% formamide	67EC; 1xSSC
L	RNA:RNA	<50	T _L *; 2xSSC	T _l *; 2xSSC
M	DNA:DNA	> 50	50EC; 4xSSC -or- 40EC; 6xSSC, 50% formamide	50EC; 2xSSC
N	DNA:DNA	<50	T _N *; 6xSSC	T _n *; 6xSSC
O	DNA:RNA	> 50	55EC; 4xSSC -or- 42EC; 6xSSC, 50% formamide	55EC; 2xSSC
P	DNA:RNA	<50	T _P *; 6xSSC	T _p *; 6xSSC
Q	RNA:RNA	> 50	60EC; 4xSSC -or- 45EC; 6xSSC, 50% formamide	60EC; 2xSSC
R	RNA:RNA	<50	T _R *; 4xSSC	T _r *; 4xSSC

¹: The hybrid length is that anticipated for the hybridized region(s) of the hybridizing polynucleotides. When hybridizing a polynucleotide to a target polynucleotide of unknown sequence, the hybrid length is assumed to be

5 that of the hybridizing polynucleotide. When polynucleotides of known

sequence are hybridized, the hybrid length can be determined by aligning the sequences of the polynucleotides and identifying the region or regions of optimal sequence complementarity.

^H: SSPE (1xSSPE is 0.15M NaCl, 10mM NaH₂PO₄, and 1.25mM EDTA, pH

- 5 7.4) can be substituted for SSC (1xSSC is 0.15M NaCl and 15mM sodium citrate) in the hybridization and wash buffers; washes are performed for 15 minutes after hybridization is complete.
- 10 *T_B - T_R: The hybridization temperature for hybrids anticipated to be less than 50 base pairs in length should be 5-10EC less than the melting temperature (T_m) of the hybrid, where T_m is determined according to the following equations. For hybrids less than 18 base pairs in length, T_m(EC) = 2(# of A + T bases) + 4(# of G + C bases). For hybrids between 18 and 49 base pairs in length, T_m(EC) = 81.5 + 16.6(log₁₀[Na⁺]) + 0.41(%G + C) - (600/N), where N is the number of bases in the hybrid,
- 15 and [Na⁺] is the concentration of sodium ions in the hybridization buffer ([Na⁺] for 1xSSC = 0.165 M).

Additional examples of stringency conditions for polynucleotide hybridization are provided in Sambrook, J., E.F. Fritsch, and T. Maniatis,

- 20 1989, Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY, chapters 9 and 11, and Current Protocols in Molecular Biology, 1995, F.M. Ausubel et al., eds., John Wiley & Sons, Inc., sections 2.10 and 6.3-6.4, incorporated herein by reference.

- 25 Preferably, each such hybridizing polynucleotide has a length that is at least 25% (more preferably at least 50%, and most preferably at least 75%) of the length of the polynucleotide of the present invention to which it hybridizes, and has at least 60% sequence identity (more preferably, at least 75% identity; most preferably at least 90% or 95% identity) with the polynucleotide of the present invention to which it hybridizes, where sequence identity is determined by comparing the
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sequences of the hybridizing polynucleotides when aligned so as to maximize overlap and identity while minimizing sequence gaps.

Expression of BBP1

5 The isolated polynucleotide of the invention may be operably linked to an expression control sequence such as the pMT2 or pED expression vectors disclosed in Kaufman et al., Nucleic Acids Res. 19, 4485-4490 (1991), in order to produce the protein recombinantly. Many suitable expression control sequences are known in the art. General methods of 10 expressing recombinant proteins are also known and are exemplified in R. Kaufman, Methods in Enzymology 185, 537-566 (1990). As defined herein "operably linked" means that the isolated polynucleotide of the invention and an expression control sequence are situated within a vector or cell in such a way that the protein is expressed by a host cell which 15 has been transformed (transfected) with the ligated polynucleotide/expression control sequence.

Expression Systems for BBP1

20 A number of types of cells may act as suitable host cells for expression of the protein. Mammalian host cells include, for example, monkey COS cells, Chinese Hamster Ovary (CHO) cells, human kidney 293 cells, human epidermal A431 cells, human Colo205 cells, 3T3 cells, CV-1 cells, other transformed primate cell lines, normal diploid cells, cell strains derived from in vitro culture of primary tissue, primary explants, 25 HeLa cells, mouse L cells, BHK, HL-60, U937, HaK or Jurkat cells.

30 Alternatively, it may be possible to produce the protein in lower eukaryotes such as yeast or in prokaryotes such as bacteria. Potentially suitable yeast strains include *Saccharomyces cerevisiae*, *Schizosaccharomyces pombe*, *Kluyveromyces* strains, *Candida*, or any yeast strain capable of expressing heterologous proteins. Potentially suitable bacterial strains include *Escherichia coli*, *Bacillus subtilis*,

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Salmonella typhimurium, or any bacterial strain capable of expressing heterologous proteins. If the protein is made in yeast or bacteria, it may be necessary to modify the protein produced therein, for example by phosphorylation or glycosylation of the appropriate sites, in order to

5 obtain the functional protein. Such covalent attachments may be accomplished using known chemical or enzymatic methods.

The protein may also be produced by operably linking the isolated polynucleotide of the invention to suitable control sequences in one or more insect expression vectors, and employing an insect expression

10 system. Materials and methods for baculovirus/insect cell expression systems are commercially available in kit form from, e.g., Invitrogen, San Diego, California, U.S.A. (the MaxBac7 kit), and such methods are well known in the art, as described in Summers and Smith, Texas Agricultural Experiment Station Bulletin No. 1555 (1987), incorporated herein by reference. As used herein, an insect cell capable of expressing a polynucleotide of the present invention is "transformed."

The protein of the invention may be prepared by culturing transformed host cells under culture conditions suitable to express the recombinant protein. The resulting expressed protein may then be

20 purified from such culture (i.e., from culture medium or cell extracts) using known purification processes, such as gel filtration and ion exchange chromatography. The purification of the protein may also include an affinity column containing agents which will bind to the protein; one or more column steps over such affinity resins as

25 concanavalin A-agarose, heparin-toyopearl7 or Cibacrom blue 3GA Sepharose7; one or more steps involving hydrophobic interaction chromatography using such resins as phenyl ether, butyl ether, or propyl ether; or immunoaffinity chromatography.

30 Alternatively, the protein of the invention may also be expressed in a form which will facilitate purification. For example, it may be expressed as a fusion protein, such as those of maltose binding protein (MBP),

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glutathione-S-transferase (GST) or thioredoxin (TRX). Kits for expression and purification of such fusion proteins are commercially available from New England BioLab (Beverly, MA), Pharmacia (Piscataway, NJ) and InVitrogen, respectively. The protein can also be tagged with an epitope

5 and subsequently purified by using a specific antibody directed to such epitope. One such epitope ("Flag") is commercially available from Kodak (New Haven, CT).

Finally, one or more reverse-phase high performance liquid chromatography (RP-HPLC) steps employing hydrophobic RP-HPLC media,

10 e.g., silica gel having pendant methyl or other aliphatic groups, can be employed to further purify the protein. Some or all of the foregoing purification steps, in various combinations, can also be employed to provide a substantially homogeneous isolated recombinant protein. The protein thus purified is substantially free of other mammalian proteins and

15 is defined in accordance with the present invention as an "isolated protein."

The protein of the invention may also be expressed as a product of transgenic animals, e.g., as a component of the milk of transgenic cows, goats, pigs, or sheep which are characterized by somatic or germ cells

20 containing a nucleotide sequence encoding the protein.

The protein may also be produced by known conventional chemical synthesis. Methods for constructing the proteins of the present invention by synthetic means are known to those skilled in the art. The synthetically-constructed protein sequences, by virtue of sharing primary,

25 secondary or tertiary structural and/or conformational characteristics with proteins may possess biological properties in common therewith, including protein activity. Thus, they may be employed as biologically active or immunological substitutes for natural, purified proteins in screening of therapeutic compounds and in immunological processes for the

30 development of antibodies.

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The proteins provided herein also include proteins characterized by amino acid sequences similar to those of purified proteins but into which modification are naturally provided or deliberately engineered. For example, modifications in the peptide or DNA sequences can be made by

- 5 those skilled in the art using known techniques. Modifications of interest in the protein sequences may include the alteration, substitution, replacement, insertion or deletion of a selected amino acid residue in the coding sequence. For example, one or more of the cysteine residues may be deleted or replaced with another amino acid to alter the conformation of the molecule. Techniques for such alteration, substitution, replacement, insertion or deletion are well known to those skilled in the art (see, e.g., USP No. 4,518,584). Preferably, such alteration, substitution, replacement, insertion or deletion retains the desired activity of the protein.
- 10 Other fragments and derivatives of the sequences of proteins which would be expected to retain protein activity in whole or in part and may thus be useful for screening or other immunological methodologies may also be easily made by those skilled in the art given the disclosures herein. Such modifications are believed to be encompassed by the present invention.
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Yeast 2 Hybrid Assays

Y2H assays demonstrated that the association of BAP with the BBP1 fusion protein is specific. The association of BBP1 with BAP suggests that BBP1 activity may have a defined role in the pathogenesis of Alzheimer's disease.

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BBP1 sequences were compared to Genbank using the basic local alignment search tool (BLAST; Altschul et al., 1990). The BBP1 protein and translations of available expressed sequence tags were aligned, searched for conserved segments, and evaluated by the MoST (Tatusov et al., 1994) protein motif search algorithm. These analyses revealed a

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potential evolutionary relationship to the G protein-coupled receptor (GPCR) family. Specifically, these analyses indicated that BBP1 contains two potential transmembrane (tm) domains equivalent to tm domains 3 and 4 of G protein-coupled receptors. The intervening hydrophilic loop

5 contains a well-characterized three amino acid motif, aspartate (D) or glutamate followed by arginine (R) and an aromatic residue (Y or F) (commonly referred to as the DRY sequence), that is conserved in almost all members of this receptor family and has been shown to serve as a molecular trigger for G protein activation (Acharya and Karnik, 1996).

10 Data from Y2H assays (see Figures 2-4) indicate that BBP1 represents a novel protein potentially containing a functional module shared with members of the G protein-coupled receptor superfamily. Specifically, it appears that BBP1 retains the critical DRF sequence (amino acids 199 to amino acids 201 of SEQ ID NO: 2), between two predicted

15 tm domains, and may have the potential to couple to a G protein regulated signaling pathway.

APP has been shown to functionally associate with G α o (Nishimoto et al., 1993; Yamatsuji et al., 1996) and BBP1 contains a structural motif known to be a G α protein activating sequence in the

20 related G protein-coupled receptors. Additionally, a hypothesis based on the predicted position and orientation of BBP1 tm domains suggests that the region of the protein that interacts with BAP would be topographically constrained to the same location as BAP in APP.

Y2H assay strains were engineered to evaluate the association of

25 the BBP1 intracellular region with G α proteins. The predicted intracellular sequences of BBP1 were expressed as a fusion protein and assayed for interactions with C-terminal regions of three G α proteins. Protein segments used in these experiments are listed in Table 2, below. The BBP1 intracellular loop interacted with all three G α proteins (Figure 3),

30 supporting the premise that BBP1 may function as a modulator of G protein activity. These various Y2H assays suggest the intriguing model

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of a multiple protein complex minimally composed of the integral membrane proteins BBP1 and APP coupled to a heterotrimeric G protein.

Table 2. Plasmids used in yeast 2-hybrid assays

expression plasmid	protein	segment
	BAP	
pEK162	(human)	1 - 42
pEK240	(mouse)	1 - 42
	BBP1	
pEK196	(clone 14)	68 - 269
pEK198	(Δtm)	68 - 202
pEK219	(ΔC)	68 - 175
pEK216	(ΔN)	123 - 202
pOZ339	(intracellular)	185 - 217
	G α	
pOZ345	(G α s)	235 - 394
pOZ346	(G α o)	161 - 302
pOZ348	(G α i2)	213 - 355

5 Further analysis of BBP1 was obtained using Y2H assays. Two overlapping portions of the BBP1 sequences contained in the BBP1Δtm clone were amplified and cloned into the Y2H vector pACT2 (expression plasmids pEK216 and pEK219, Table 2 and corresponding proteins BBP1ΔN and BBP1ΔC, (Figure 4)). The ΔC construct lacked both tm
10 domains; the ΔN construct encoded the first tm domain plus the preceding 52 amino acids. These fusion proteins were assayed with the BAP fusion protein and responses compared to those of strains expressing the larger BBP1Δtm protein. The BBP1ΔC protein induced a weak Y2H response (compare BBP1ΔC to vector, Figure 4), but the BBP1ΔN protein,
15 containing the first tm domain and adjacent amino-proximal sequences produced a response only slightly weaker than that observed with BBP1Δtm (Figure 4). These results suggest that a major determinant for

the association with BAP is contained within the BBP1 region predicted to be topographically similar to BAP in the wild-type APP protein.

The Y2H system was utilized to demonstrate the selectivity and specificity of BBP1 binding to human BAP as compared to rodent BAP.

5 There are three amino acid substitutions (G5R, F10Y and R13H) in the rodent BAP sequence compared to the human sequence. It was of interest, to evaluate the association of rodent BAP with BBP1 in the Y2H system. The sequence of human BAP in pEK162 was changed to encode the rodent peptide by oligonucleotide directed mutagenesis by PCR. The 10 resultant plasmid, pEK240, is identical to the human BAP fusion protein expression plasmid utilized throughout this report except for the three codons producing the amino acid substitutions for the rodent peptide sequence. Interactions between BBP1 fusion protein and rodent and human BAP fusion proteins were compared by Y2H bioassay. Strains 15 expressing BBP1 and the rodent BAP failed to produce a growth response (Figure 7). This finding supports the premise that BBP1 may serve as a specific mediator of the neurotoxic effects of BAP, and provides a mechanism to explain the reduced neurotoxicity of the rodent BAP. Importantly, these data also serve to illustrate the high degree of 20 specificity of the BBP1/BAP interaction in the Y2H assays since the substitution of three amino acids was sufficient to completely abrogate the association (Figure 7).

Isolated BBP1 polypeptides

25 Proteins and protein fragments of the present invention include proteins with amino acid sequence lengths that are at least 25% (more preferably at least 50%, and most preferably at least 75%) of the length of a disclosed protein and have at least 60% sequence identity (more preferably, at least 75% identity; most preferably at least 90% or 95% 30 identity) with that disclosed protein, where sequence identity is determined by comparing the amino acid sequences of the proteins when

aligned so as to maximize overlap and identity while minimizing sequence gaps. Also included in the present invention are proteins and protein fragments that contain a segment preferably comprising 8 or more (more preferably 20 or more, most preferably 30 or more) contiguous amino

5 acids that shares at least 75% sequence identity (more preferably, at least 85% identity; most preferably at least 95% identity) with any such segment of any of the disclosed proteins.

Species homologues of the disclosed polynucleotides and proteins are also provided by the present invention. As used herein, a species

10 homologue is a protein or polynucleotide with a different species of origin from that of a given protein or polynucleotide, but with significant sequence similarity to the given protein or polynucleotide. Preferably, polynucleotide species homologues have at least 60% sequence identity (more preferably, at least 75% identity; most preferably at least 90% 15 identity) with the given polynucleotide, and protein species homologues have at least 30% sequence identity (more preferably, at least 45% identity; most preferably at least 60% identity) with the given protein, where sequence identity is determined by comparing the nucleotide sequences of the polynucleotides or the amino acid sequences of the 20 proteins when aligned so as to maximize overlap and identity while minimizing sequence gaps. Species homologues may be isolated and identified by making suitable probes or primers from the sequences provided herein and screening a suitable nucleic acid source from the desired species. Preferably, species homologues are those isolated from 25 mammalian species. Most preferably, species homologues are those isolated from certain mammalian species such as, for example, *Pan troglodytes*, *Gorilla gorilla*, *Pongo pygmaeus*, *Hylobates concolor*, *Macaca mulatta*, *Papio papio*, *Papio hamadryas*, *Cercopithecus aethiops*, *Cebus capucinus*, *Aotus trivirgatus*, *Sanguinus oedipus*, *Microcebus murinus*, 30 *Mus musculus*, *Rattus norvegicus*, *Cricetulus griseus*, *Felis catus*, *Mustela vison*, *Canis familiaris*, *Oryctolagus cuniculus*, *Bos taurus*, *Ovis aries*, *Sus*

scrofa, and Equus caballus, for which genetic maps have been created allowing the identification of syntenic relationships between the genomic organization of genes in one species and the genomic organization of the related genes in another species (O'Brien and Seuanez, 1988, Ann. Rev.

5 Genet. 22: 323-351; O'Brien et al., 1993, Nature Genetics 3:103-112; Johansson et al., 1995, Genomics 25: 682-690; Lyons et al., 1997, Nature Genetics 15: 47-56; O'Brien et al., 1997, Trends in Genetics 13(10): 393-399; Carver and Stubbs, 1997, Genome Research 7:1123-1137; all of which are incorporated by reference herein).

10 The invention also encompasses allelic variants of the disclosed polynucleotides or proteins; that is, naturally-occurring alternative forms of the isolated polynucleotides which also encode proteins which are identical or have significantly similar sequences to those encoded by the disclosed polynucleotides. Preferably, allelic variants have at least 60%
15 sequence identity (more preferably, at least 75% identity; most preferably at least 90% identity) with the given polynucleotide, where sequence identity is determined by comparing the nucleotide sequences of the polynucleotides when aligned so as to maximize overlap and identity while minimizing sequence gaps. Allelic variants may be isolated and identified
20 by making suitable probes or primers from the sequences provided herein and screening a suitable nucleic acid source from individuals of the appropriate species.

The invention also includes polynucleotides with sequences complementary to those of the polynucleotides disclosed herein.

25

Applications

BBP1 proteins of the present invention can be used in a variety of applications routine to one of skill in the art based upon this disclosure.

Specifically the BBPs can be used as immunogens to raise antibodies

30 which are specific to the cloned polypeptides. Various procedures known in the art may be used for the production of antibodies to BBP1 proteins.

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Such antibodies include, but are not limited to, polyclonal, monoclonal, chimeric, single chain, Fab fragments and an Fab expression library. For the production of antibodies, various host animals including, but not limited to rabbits, mice, and rats, are injected with a BBP. In one

5 embodiment, the polypeptide or a fragment of the polypeptide capable of specific immunoactivity is conjugated to an immunogenic carrier.

Adjuvants may also be administered in conjunction with the polypeptide to increase the immunologic response of the host animal. Examples of adjuvants which may be used include, but are not limited to, complete

10 and incomplete Freund's, mineral gels such as aluminum hydroxide, surface active substances such as lysolecithin, pluronic polyols, polyanions, peptides, oil emulsions, keyhole limpet hemocyanin, and dinitrophenol.

Monoclonal antibodies to BBP1 proteins of the present invention

15 can be prepared using any technique which provides for the production of antibodies by continuous cell line in culture. Such techniques are well known to those of skill in the art and include, but are not limited to, the hybridoma technology originally described by Kohler and Milstein (Nature 1975, 256,4202-497), the human B-cell hybridoma technique described
20 by Kosbor et al. (Immunology Today 1983, 4, 72) and the EBV-hybridoma technique described by Cole et al. (Monoclonal Antibodies and Cancer Therapy, Alan R. Liss, Inc., pp 77-96).

Antibodies immunoreactive to the polypeptides of the present invention can then be used to screen for the presence and subcellular
25 distribution of similar polypeptides in biological samples. In addition, monoclonal antibodies specific to the BBP1 proteins of the present invention can be used as therapeutics.

The BBP1 proteins can also serve as antigens useful in solid phase assays measuring the presence of antibodies which immunoreact with the
30 claimed peptides. Solid phase competition assays can be used to measure immunological quantities of clone 14-related antigen in biological

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samples. This determination is not only useful in facilitating the complete characterization of the cellular function or functions of the polypeptides of the present inventions, but can also be used to identify patients with abnormal amounts of these proteins.

5 BBP1 proteins of the present invention can also be used as capture reagents in affinity chromatography for the detection of BAP and BAP aggregates as markers for AD.

In addition, these BBP1s are useful as reagents in an assay to identify candidate molecules which effect the interaction of BAP and the 10 cloned protein. Compounds that specifically block this association could be useful in the treatment or prevention of AD.

These BBP1s are also useful in acellular in vitro binding assays wherein alteration by a compound in the binding of these beta amyloid peptide associated proteins to BAP or BAP aggregates is determined.

15 Acellular assays are extremely useful in screening sizable numbers of compounds since these assays are cost effective and easier to perform than assays employing living cells. Upon disclosure of the polypeptides of the present invention, the development of these assays would be routine to the skilled artisan. In such assays, either BBP1 or BAP is labeled.

20 Such labels include, but are not limited to, radiolabels, antibodies, and fluorescent or ultraviolet tags. Binding of a BBP1 to BAP or BAP aggregates is first determined in the absence of any test compound. Compounds to be tested are then added to the assay to determine whether such compounds alter this interaction.

25

Examples

The present invention is further described by the following examples. The examples are provided solely to illustrate the invention by reference to specific embodiments. These exemplifications, while 30 illustrating certain specific aspects of the invention do not portray the limitations or circumscribe the scope of the invention.

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Yeast two-hybrid system (hereinafter "Y2H"): Y2H expression plasmids were constructed in vectors pAS2 and pACT2 (described in WadeHarper et al., 1993) and pCUP (described in Ozenberger and Young, 1995). Yeast strain CY770 (Ozenberger and Young, 1995) served as the host for all Y2H assays.

Genetic screen: The polymerase chain reaction (PCR) method was used to amplify and modify sequences encoding BAP. Oligonucleotides #1 (5' - CC ATG GAT GCA GAA TTC CGA C) and #3 (5' - AAGCTTGTGAC TTA CGC TATGAC AAC ACC GC) were used to amplify BAP using pCLL621, a modified human APP clone (Jacobsen et al., 1994), as template. The amplified DNA consists of codons 389 to 430 (which encodes BAP₄₂) of the APP precursor protein with the following modifications. The sense strand primer added a 5' Ncol restriction site in the same translational reading frame as the Ncol site in pAS2. The antisense strand primer added a stop codon and HindIII and Sall sites for cloning. The product from this amplification was ligated into the TA cloning system (Invitrogen Corp., Carlsbad, CA), and subsequently removed by digestion with Ncol and Sall. This fragment was cloned into pAS2 cleaved with Ncol plus Sall. The resultant plasmid, pEK162, was confirmed by DNA sequencing through the GAL4/BAP junction. The protein (BAP^{BD}; Figure 1) expressed from pEK162 comprised a fusion protein containing the DNA-binding domain of the yeast transcriptional activation protein Gal4 (lacking functional activation sequences) with the addition of the 42 amino acids of BAP to the carboxy-terminus. An expression plasmid was developed that mediates the expression of unmodified BAP₄₂. Oligo #2 (5' - AAGCTTAAG ATG GAT GCA GAA TTC CGA C) was paired with oligo #3 in a PCR as described above. The product of this amplification contains a 5' HindIII site and translation initiation signals optimized for expression in *Saccharomyces cerevisiae*. Again, the DNA fragment was cloned into the TA system. It was then isolated on a HindIII fragment and cloned into pCUP cleaved with HindIII.

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The orientation of the BAP gene in the resultant plasmid, pEK149 (BAP; Figure 1), was confirmed by DNA sequencing. The BAP expression plasmids pEK149 (which used URA3 as the selection marker) and pEK162 (which used TRP1 as the selection marker) were transformed into the 5 yeast host CY770 (Ozenberger and Young, 1995). The strain containing both plasmids was designated CY2091. A plasmid library consisting of cDNA fragments isolated from human fetal brain cloned into the yeast 2-hybrid expression vector pACT2 (which used LEU2 as the selection marker) was purchased from Clontech Laboratories, Inc. (Palo Alto, CA).

10 The library-derived protein is depicted in Figure 1 as unknown^{AD}. This library was used to transform CY2091. The samples were spread on synthetic complete (SC) yeast growth medium lacking uracil, tryptophan, and leucine to select cells containing all three plasmids. The medium also lacked histidine and contained 3-amino-triazole, an inhibitor of the product

15 of the yeast HIS3 gene, at a concentration of 25 mM. 3-Amino-triazole was utilized to reduce activity from low-level constitutive expression of the HIS3 reporter gene. Plates were incubated at 30°C for 12 days. Twenty-four colonies exhibiting increased histidine prototrophy were isolated. Transformation controls indicated that the screen assayed 10⁶

20 individual clones. A PCR approach was utilized to quickly determine the content of positive clones. Total DNA was isolated from each positive strain by standard methods. This material was used as template for PCRs using oligos #4 (5' - TTTAATACCA CTACAATGGA T) plus #5 (5' - TTTTCAGTAT CTACGATTCA T) which flank the cloning region of the

25 library vector pACT2. DNA fragments were ligated into the TA system and examined by DNA sequencing. The library plasmid contained in clone #14 (as described above) was isolated by shuttle into E. coli. The nucleotide sequence of the human cDNA sequences was determined, confirming the sequence of the initial PCR product.

30 Bioassays: Strains were grown overnight in 2ml SC medium lacking leucine and tryptophan to a density of approximately 7 x 10⁷ cells

per ml. Cells were counted and 10-fold serial dilutions made from 10^4 to 10^8 cells per ml in sterile water. These samples were spotted in 5 μ l aliquots on SC medium lacking leucine, tryptophan and histidine and containing 25 mM 3-amino-triazole. Plates were incubated at 30°C for 2 to 3 days. Positive protein/protein interactions were identified by increased prototrophic growth compared to control strains expressing the Gal4 DNA-binding domain fusion protein plus an irrelevant transcriptional activation domain fusion protein (or simply containing the pACT vector without inserted sequences). These control strains were indicated in the Figures described above as the label 'vector'. This assay method was highly reproducible and provided for the detection of subtle inductions of growth mediated by the specific interaction between target proteins. The original BBP1 clone, designated pEK196 and deposited as ATCC 98399; is referred herein as clone 14), was used as a PCR template to truncate the protein product to express BBP1 Δ tm. Sense primer #6 (5'-TTTAATACCA CTACAATGGA T) annealed to GAL4 sequences in pACT2. The antisense primer #7 (5'-CTCGAG TTA AAA TCG ATC TGC TCC CAA CC) incorporated a 3' stop codon and Xhol site immediately 3' to the sequences encoding the DRF motif of BBP1. The PCR product was ligated into the TA cloning vector and subsequently digested with EcoRI + Xhol and cloned into pACT2. The hybrid product expressed from this plasmid (pEK198) was denoted BBP1 Δ tm. Similarly, primer #7 was paired with primer #8 (5'-GAATT CCA AAA ATA AAT GAC GCT ACG) to engineer the BBP1 Δ N expression plasmid pEK216. Again, the PCR product was ligated into the TA system and the resultant plasmid digested with EcoRI + Xhol with the BBP1 fragment (codons 123-202) finally ligated into pACT2 digested with the same enzymes. BBP1 Δ C was made by using the pACT2-specific oligo #6 with antisense oligo #9 (5'-CTCGAG TCA AGA TAT GGG CTT GAA AAA AC). After TA cloning, isolation of the EcoRI-Xhol fragment and cloning into pACT2, the resultant plasmid, pEK219, expressed BBP1 from residue 68 to 175. Sequences

2025262728292A2B2C2D2E2F2G2H2I2J2K2L2M2N2O2P2Q2R2S2T2U2V2W2X2Y2Z2

encoding the BBP1 intracellular loop were amplified using oligonucleotides #10 (5'-CCTTCC ATG GAA GTG GCA GTC GCA TTG TCT) plus #11 (5'-AACACTCGAG TCA AAA CCC TAC AGT GCA AAA C). This product, containing BBP1 codons 185 to 217, was digested with Ncol + Xhol and

5 cloned into pAS2 cleaved with Ncol + Sall to generate pOZ339.

Construction of all G α protein expression plasmids utilized the BamHI site near the center of each rat cDNA sequence (Kang et al., 1990) as the site of fusion in pACT2. Sense primers annealed to sequences 5' of the BamHI site; antisense primers annealed to sequences 3' of the stop codon

10 and included a Sall restriction site. Primers were: Gao, sense (#17) = 5'-GTGGATCCAC TGCTTCGAGG AT, antisense (#18) = 5'-GTCGACGGTT GCTATACAGG ACAAGAGG; Gas, sense (#19) = 5'-GTGGATCCAG TGCTTCAATG AT, antisense (#20) = 5'-GTCGACTAAA TTTGGCGTT CCCTTCTT; Gai2, sense (#21) = 5'-GTGGATCCAC

15 TGCTTGAGG GT, antisense (#22) = 5'-GTCGACGGTC TTCTTGCCCC CATCTTCC. PCR products were cloned into the TA vector. G α sequences were isolated as BamHI-Sall fragments and cloned into pACT2 digested with BamHI + Sall. See Table 2 for plasmid designations.

Finally, oligonucleotide #23 was synthesized for the conversion of human

20 BAP to the rodent sequence. This primer has the sequence 5'-ATATGCCATG GAT GCA GAA TTC GGA CAT GAC TCA GGA TT GAA GTT CGT. The triplets represent the first 13 codons of BAP; the three nucleotides that were changed to produce the rodent sequence are underlined. Oligo #23 was paired with #24 (5'-TGACCTACAG

25 GAAAGAGTTA) which anneals to a region of the Y2H vectors that is 3' of the cloning site in a PCR using pEK162 as the template. The product was cleaved with Ncol + Sall and ligated into pAS2 to produce pEK240. The nucleotide sequence of the segment encoding rodent BAP was confirmed.

30 Genomic cloning; RACE (rapid amplification of cDNA ends): A human genomic lambda library (Stratagene), corresponding to \approx 2.0 X 10⁶ pfus,

was screened with randomly-primed EcoRI/Clal fragment probe corresponding to nucleotides 187-600 (Figure 2). The probe was labeled with [³²P]-CTP using the ⁷⁷QuickPrimer Kit according to the manufacturer's (Pharmacia) protocol. Filters were hybridized under high

5 stringency: 40°C in 50% formamide, 0.12M NaHPO₄, 0.25M NaCl, 7% SDS and 25mg/ml sonicated salmon sperm DNA and washed at 65°C in 0.1X SSC containing 0.1% sodium dodecyl sulfate and exposed to Kodak BioMax MS film. Lambda phage clones hybridizing to the probe were plaque purified by successive plating and rescreening. Ten positive clones
10 were purified and subjected to further analysis by hybridization to a 45 nt oligonucleotide probe directed to the most 5' sequences known from the original cDNA clone. This oligonucleotide was the reverse complement of nucleotides 157-201 (Figure 2) and has the sequence 5'-CCAGGCAGGCC
GCCATCTTGG AGACCGACAC TTTCTCGCCA CTTCC. Lambda phage
15 DNA was isolated by standard molecular biology techniques and subjected to direct sequencing using fluorescent dideoxy cycle sequencing on an ABI 373 sequencer.

RACE: First strand DNA synthesis was performed using the rTth thermal-stable polymerase system (Perkin Elmer). The following reagents
20 were combined in a 1.5 mL tube to give a 10 microliter volume: 1X reverse transcription buffer, 1 mM MnCl₂, 1.6 mM dNTP mix, 2.5U rTth polymerase, 100ng human hippocampus poly A⁺ RNA (Clontech), 10mM oligonucleotide (nt 429-452, Figure 2; 5'-GTTATGTTGG GTGCTGGAAA
ACAG). The reaction was incubated at 70°C for 15 minutes and
25 immediately placed on ice. The Marathon cDNA synthesis kit (Clontech) was used for second strand cDNA generation. The entire 10 μ l from the first strand reaction was combined with the following reagents: 1X second strand buffer, 0.8 mM dNTP mix, 4X second strand cocktail (E.coli DNA polymerase I, E.coli DNA ligase, E.coli RNaseH), and dH₂O up
30 to a volume of 80 μ l. The tube was incubated at 16°C for 1.5 hours after which time T4 DNA polymerase (10U) was added and incubated for an

additional 45 minutes at 16°C. To terminate the reaction, 4μl of 20X EDTA/glycogen (0.2M EDTA/2mg/ml glycogen) was added to the reaction mixes followed by a phenol/chloroform/isoamyl alcohol extraction to remove enzymes and other impurities. The DNA was precipitated by

- 5 adding 0.1X volume 3M Na acetate pH 5.2 and 2.5X volume reagent grade EtOH and place at -70°C. The DNA was washed once with 70% EtOH, dried down and resuspended in 10μl dH₂O. Half of the DNA was used for Marathon adaptor ligation to be used in subsequent RACE PCR reactions following the Clontech protocol as follows: 5μl cDNA was
- 10 added to 2μl (10mM) Marathon (5'- CTAATACGAC TCACTATAGG GCTCGAGCGG CCGCCCGGGC AGGT), 1X DNA ligation buffer and 1μl (1U) T4 DNA ligase. The reaction mix was incubated overnight at 16°C. The mix was diluted 1:50 for initial RACE reaction and combined in a 0.2mL PCR tube with the following: 40μl dH₂O, 1μl 10X KlenTaq DNA
- 15 polymerase (Clontech), 1μl (10mM) AP1 primer (5'-CCATCCTAAT ACGACTCACT ATAGGGC), 1μl (10mM) BBP1-specific primer (corresponding to nts. 187-209, Figure 2; 5'-CCAGACGGCCA GGCAGGCC AT), 5μl 10X KlenTaq polymerase buffer, 1μl 10mM dNTP mix, 1μl of diluted cDNA from above reaction. The following cycling
- 20 conditions were performed using a Perkin Elmer GeneAmp PCR system 2400 thermocycler: Denaturing cycle 94°C for 1 minute followed by 5 cycles of 30" at 94°C, 3' at 72°C, 5 cycles of 30" at 94°C, 3' at 70°C, followed by 25 cycles of 30" at 94°C, 3' at 68°C, with a final extension 7' at 72°C. This was followed by a nested RACE PCR reaction as
- 25 follows: 40μl dH₂O, 1μl (1U) AmpliTaqGold DNA polymerase (Perkin Elmer), 1μl (10mM) AP2 primer (5'-ACTCACTATA GGGCTCGAGC GGC), 1μl (10mM) BBP1-specific primer (corresponding to nts. 172-194, Figure 2; 5'-GCCGCCATCT TGGAGACCGA CAC), 5μl 10X AmpliTaq polymerase buffer, 1μl 10mM dNTP mix, 1μl of primary RACE product. The PCR

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cycling conditions were an initial denaturing cycle of 9' at 94°C, 25 cycles of 30" at 94°C, 30" at 68°C, 2' at 72°C, followed by a 72°C extension for 7'. The PCR product was run on a 1% agarose gel in 1XTBE buffer. The resulting 350 base pairs product was gel purified and directly cloned using the TA Cloning Kit (Invitrogen). Ligation mixes were transformed into OneShot Cells (Invitrogen) and plated on LB-ampicillin (100 μ g/ml) agar plates containing X-gal. Mini-prep DNA was obtained and examined by fluorescent dideoxy cycle sequencing on an ABI 373 sequencer.

10 Northern analyses. Human multiple tissue and multiple brain tissue mRNA Northern blots were obtained from Clontech (Palo Alto, CA). BBP1 sequences extending from the original fusion junction to the poly-A region were isolated on an EcoRI fragment from a TA clone derived from pEK196. β -actin DNA was provided by the manufacturer. Radiolabelled probes were produced from these DNAs using a random priming method to incorporate 32 P-dCTP (Pharmacia Biotech, Piscataway, NJ).

15 Hybridizations were performed per manufacturer's (Clontech) instructions in Express Hyb Solution at 68°C. Blots were washed in 2x SSC (1X SSC is 0.15 M sodium chloride, 0.015 M sodium citrate), 0.05% SDS at room temperature, followed by two washes in 0.1 x SSC, 0.1% SDS at 50°C.

20 Hybridization signals were visualized by exposure to Kodak BioMax film.

In situ hybridization. DNA templates for riboprobe synthesis were prepared by PCR using a plasmid clone containing the full length human BBP cDNA. A single riboprobe targeted to the 3' UTR of the cDNA was used. The probe sequences were checked versus the GenBank database to ensure that they only recognize the appropriate targets among all deposited sequences. To generate riboprobes for BBP1, a pair of oligonucleotide primers was designed to amplify a 275 base pairs region from the 3' UTR of the BBP1 cDNA and, in addition, add the promoter sequences for T7 (sense) and T3 (antisense) polymerase. These primers contained the following sequences: 5'-TAATACGACT CACTATAGGG

TTAGAAGAAA CAGATTGAG (forward); 5'-ATTAACCCTC
ACTAAAGGGA CAAGTGGCAA CTTGCCTTG (reverse). PCR products
were gel purified on 1.5% low-melt agarose gels, and bands containing
the products were excised, phenol and phenol-chloroform extracted, and

5 ethanol precipitated. Pellet were dried and resuspended in 1X TE buffer
(10 mM Tris-HCl, 1mM EDTA, pH 7.4). The APP riboprobe template
consisted of a Ddel-Xhol fragment from the protein coding region, as
described by Jacobsen et al. (1991). Fifty ng of DNA template was used
for transcription reactions using (³⁵S)-CTP (New England Nuclear, Boston,
10 MA) and the Riboprobe Gemini™ System (Promega, Madison, WI).

In situ hybridization histochemistry using sections of postmortem
human hippocampus were performed as described previously (Rhodes,
1996). Sections were cut at 10 µm on a Hacker-Brights cryostat and
thaw-mounted onto chilled (-20°C) slides coated with Vectabond reagent

15 (Vector Labs, Burlingame, CA). All solutions were prepared in dH₂O
treated with 0.1% (v/v) diethylpyrocarbonate and autoclaved. Sections
were fixed by immersion in 4% paraformaldehyde in PBS (pH 7.4) then
immersed sequentially in 2xSSC, dH₂O, and 0.1M triethanolamine, pH
8.0. The sections were then acetylated by immersion in 0.1M
20 triethanolamine containing 0.25% (v/v) acetic anhydride, washed in
0.2xSSC, dehydrated in 50, 70 and 90% ethanol, and rapidly dried. One
ml of prehybridization solution containing 0.9M NaCl, 1mM EDTA, 5x
Denhardt's, 0.25 mg/ml single-stranded herring sperm DNA (GIBCO/BRL,
Gaithersberg, MD), 50% deionized formamide (EM Sciences, Gibbstown,
25 NJ) in 10mM Tris, (pH 7.6), was pipetted onto each slide, and the slides
incubated for 3 hrs. at 50°C in a humidified box. The sections were then
dehydrated by immersion in 50, 70, and 90% ethanol and air dried.
Labeled riboprobes were added at a final concentration of 50,000 cpm/µl
30 to hybridization solution containing 0.9M NaCl, 1mM EDTA, 1x
Denhardt's, 0.1 mg/ml yeast tRNA, 0.1 mg/ml single-stranded salmon
sperm DNA, dextran sulfate (10%), 0.08% BSA, 10mM DTT (Boehringer

Mannheim, Indianapolis, IN), and 50% deionized formamide in 10mM Tris (pH 7.6). The probes were then denatured at 95°C (1 min), placed on ice (5 min), and pipetted onto the sections and allowed to hybridize overnight at 55°C in a humidified chamber. The sections were subsequently

5 washed 1 x 45 min at 37°C in 2xSSC containing 10mM DTT, followed by 1 x 30 min at 37°C in 1xSSC containing 50% formamide, and 1 x 30 min at 37°C in 2xSSC. Single stranded and non-specifically hybridized riboprobe was digested by immersion in 10mM Tris pH 8.0 containing bovine pancreas RNase A (Boehringer Mannheim; 40 mg/ml), 0.5M NaCl, 10 and 1mM EDTA. The sections were washed in 2XSSC for 1 hr at 60°C, followed by 0.1XSSC containing 0.5% (w/v) sodium thiosulfate for 2 hrs. at 60°C. The sections were then dehydrated in 50, 70, 90% ethanol containing 0.3M ammonium acetate, and dried. The slides were loaded into X-ray cassettes and opposed to Hyperfilm b-Max (Amersham) for 14- 15 30 days. Once a satisfactory exposure was obtained, the slides were coated with nuclear-track emulsion (NTB-2; Kodak) and exposed for 7-21 days at 4°C. The emulsion autoradiograms were developed and fixed according to the manufacturer's instructions, and the underlying tissue sections were stained with hematoxylin. To assess nonspecific labeling, a 20 control probe was generated from a template provided in the Riboprobe Gemini™ System kit (Promega). This vector was linearized using Scal and transcribed using T3 polymerase. The resulting transcription reaction generates two products, a 250 base and a 1,525 base riboprobe, containing only vector sequence. This control probe mixture was labeled 25 as described above and added to the hybridization solution at a final concentration of 50,000 cpm/μl. No specific hybridization was observed in control sections, i.e., these sections gave a very weak uniform hybridization signal that did not follow neuroanatomical landmarks (data not shown).

30 Example 1: Cloning and Isolation BAP-binding protein (BBP1).

A yeast 2-hybrid (Y2H) genetic screen was developed to identify proteins which interact with human BAP₄₂, a 42 amino acid proteolytic fragment of APP which is considered to potentially be the more toxic aggregated form of BAP. BAP₄₂ was expressed fused to the yeast Gal4

5 DNA-binding domain and was also expressed as free peptide (Figure 1). This strain was transformed with a human fetal brain cDNA Y2H library. A single clone, designated clone 14 defined above, from approximately 10⁶ independent transformants, produced consistent reporter gene activation and contained a substantial open reading frame continuous with that of 10 the GAL4 domain. The cDNA insert comprised 984 base pairs, terminating in a poly-A tract. This sequence encoded 201 amino acids (SEQ ID NO: 2; amino acid residues 68 to 269) with two regions of sufficient length and hydrophobicity to transverse a cellular membrane.

The library-derived plasmid was isolated from clone 14 and used to 15 reconstruct Y2H assay strains. Examination of these strains demonstrated that the BAP fusion protein specifically interacted with the clone 14 protein, although the response was weak. Since protein domains of strong hydrophobicity, such as transmembrane regions, inhibit 20 Y2H responses (Ozenberger, unpublished data), clone 14 insert was truncated (hereinafter BBP1Δtm) to remove the region of strongest hydrophobicity and retested for interactions with BAP. A much more robust Y2H response was observed with BBP1Δtm (Figure 2), supporting the notion that the deleted sequences encode a potential transmembrane ("tm") anchor. The nucleotide sequence of Clone 14 was searched 25 against GenBank; the BAP binding protein (BBP1) thus identified appeared to be novel.

Example 2: Isolation and confirmation of the 5' terminus of BBP1.

The BBP1 cDNA sequences contained in clone 14 described in 30 Example 1, above, lacked the 5' end of the protein coding region as no potential initiating methionine codon was present. Multiple attempts at

conventional 5' RACE (rapid amplification of cDNA ends) utilizing a standard reverse-transcriptase only resulted in the addition of 27 nucleotides. These sequences included an ATG, but no upstream stop codon in the same translational reading frame to provide confidence that

5 this was the initiating codon. A genomic cloning approach was initiated to isolate the 5' terminus of the BBP1 gene.

Hybridization of a human genomic lambda library with a randomly-primed probe corresponding to 400 base pairs (bps) of the 5' sequence of clone 14 resulted in identification of 10 positive clones. Further

10 characterization of these clones using a 45-base oligonucleotide probe directed to the most upstream BBP1 sequence of clone 14 (and corresponding to the 5' upstream sequence of the 400 base pairs probe revealed that 6 of the 10 clones included the terminal 5' sequences contained within those previously identified. It was determined that the

15 other 4 lambda clones represented other exons which were contained within the original 400 base pairs randomly-primed cDNA-derived probe (data not shown). Direct cycle sequencing of lambda phage DNA from representative clones corresponding to the 5' end of BBP1 revealed ^a500 nucleotides upstream and overlapping with the sequence known for clone

20 14. This additional sequence potentially encodes 62 additional amino acids upstream of the previously characterized MET before arriving at a MET preceded by an in-frame stop codon. Although there exist two MET residues downstream from the furthest upstream MET, by standard convention we have tentatively defined the sequence of the amino

25 terminus of the human BBP1 gene to include the first 5' MET which follows an in-frame stop codon. The entire coding region and deduced protein sequence is shown in SEQ ID NOS:1 and 2. A plasmid (denoted BBP1-f1) containing this amino acid sequence has been deposited in the American Type Culture Collection having accession number 98617).

30 Since the 5' coding sequences were derived from a genomic library, there existed the possibility that this region contained introns.

This potentiality was investigated by two methods. First, a forward primer directed to the region of the 5' MET and a reverse primer within the original clone 14 were utilized to amplify sequences from brain cDNA as well as from genomic DNA. Products of identical size were generated 5 from both samples, indicating the absence of introns within this region and confirming the linkage of the upstream sequence with the original sequence. Secondly, cDNA sequences were isolated in modified 5' RACE experiments (see Materials and Methods, above) that were identical to those obtained from the genomic clone. These findings confirmed the 10 upstream sequences (both from genomic and cDNA sources) and the lack of introns in this region.

Example 3: Characterization of BBP1.

BBP1 sequences were compared to Genbank using the basic local 15 alignment search tool (BLAST; Altschul et al., 1990). Two *Caenorhabditis elegans* and one *Drosophila melanogaster* genomic sequence and a large number of human, mouse and other mammalian expressed sequence tags were identified. However, no complete cDNA sequences were available nor were any functional data attributed to the gene. The BBP1 protein 20 and translations of available expressed sequence tags were aligned, searched for conserved segments, and evaluated by the MoST (Tatusov et al., 1994) protein motif search algorithm. These analyses revealed a potential evolutionary relationship to the G protein-coupled receptor family. Specifically, these analyses indicated that BBP1 contains two 25 potential transmembrane (tm) domains equivalent to tm domains 3 and 4 of G protein-coupled receptors. The intervening hydrophilic loop contains a well-characterized three amino acid motif, aspartate (D) or glutamate followed by arginine (R) and an aromatic residue (Y or F) (commonly referred to as the DRY sequence), that is conserved in almost all members 30 of this receptor family and has been shown to serve as a molecular trigger for G protein activation (Acharya and Karnik, 1996). These data indicate

that BBP1 represents a novel protein potentially containing a functional module shared with members of the G protein-coupled receptor superfamily. Specifically, it appears that BBP1 retains the critical DRF sequence between two predicted tm domains, so may have the potential to couple to a G protein regulated signaling pathway.

Structural analysis of BBP1 indicated it contained a structural motif known to be a G α protein activating sequence in the related G protein-coupled receptors. Y2H assays demonstrating the interaction of BBP1 with various members of the G protein coupled receptors are illustrated in Figure 3. Based on structural predictions, BBP1 is depicted as transversing a membrane twice with both termini in the luminal compartment. Other orientations cannot be entirely ruled out. The potential protein interactions described above were investigated in Y2H assays. Two overlapping portions of the BBP1 sequences contained in the BBP1 Δ tm clone were amplified and cloned into the Y2H vector pACT2 (expression plasmids pEK216 and pEK219, Table 2 and corresponding proteins BBP1 Δ N and BBP1 Δ C, Figure 4). The Δ C construct is lacking both tm domains; the Δ N construct encodes the first tm domain plus the preceding 52 amino acids. These fusion proteins were assayed with the BAP fusion protein and responses compared to those of strains expressing the larger BBP1 Δ tm protein. These results suggest that a major determinant for the association with BAP is contained within the BBP1 region predicted to be topographically similar to BAP in the wild-type APP protein.

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Example 4: Tissue distribution of human BBP1 expression.

Expression of BBP1 mRNA was evaluated as an initial step in elucidating the activity of the gene and its product. A major transcript of 1.25 kb was observed in all tissues (Figure 5A). There was a high level of expression in heart. Whole brain exhibited an intermediate level of expression. Samples derived from separate brain regions all exhibited

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BBP1 expression (Figure 5B). Interestingly, limbic regions contained relatively greater amounts of BBP1 mRNA. These are the regions of the brain where BAP aggregation and associated neurotoxicity initially occur.

Analysis of *in situ* hybridization autoradiograms obtained using a BBP1-

5 specific riboprobe indicated that in human hippocampus and entorhinal cortex, BBP1 mRNA is expressed in medium to large cells in a pattern consistent with expression in neurons as opposed to glial cells (Figure 6). Moreover, BBP1 mRNA is expressed in virtually all hippocampal and entorhinal neurons, i.e., there do not appear to be any real or laminar 10 differences in the intensity of the hybridization signal. Interestingly, the pattern of BBP1 expression was strikingly similar to the pattern observed using a riboprobe directed against mRNA for the amyloid precursor protein APP (Figure. 6). In summary, BBP1 mRNA was observed in all tissues and all brain regions examined. *In situ* analysis of BBP1 mRNA expression 15 also revealed extensive expression in the hippocampus region.

Example 5: Cell line distribution of BBP1 expression.

BBP1 expression was also investigated in numerous cell lines and data were extracted from dbEST, the collection of expressed sequence

20 tags from the National Center for Biotechnology Information. Reverse-transcription polymerase chain reaction (RT-PCR) methods were utilized to qualitatively assess BBP1 mRNA expression in cell lines commonly utilized for recombinant protein expression as well as a variety of cancer cell lines. BBP1 was observed in hamster CHO and human HEK293 cells.

25 Signals were observed in the embryonic stem cell line Ntera-2 and neuroblastoma lines IMR32 and SK-N-SH. BBP1 expression was observed in cancer cell lines representing the following tissue origins: colon (Cx-1, Colo205, MIP101, SW948, CaCo, SW620, LS174T), ovarian (A2780S, A2780DDP), breast (MCF-7, SKBr-3, T47-D, B7474), lung (Lx-1, A5439), 30 melanoma (Lox, Skmel30), leukemia (HL60, CEM), prostate (LNCAP, Du145, PC-3). A Northern blot probing mRNA isolated from the following

cancer cell lines demonstrated BBP1 expression in all samples: promyelocytic leukemia (HL-60), carcinoma (HeLa S3), chronic myelogenous leukemia (K-562), lymphoblastic leukemia (MOLT-4), Burkitt's lymphoma (Raji), colorectal adenocarcinoma (SW480), lung carcinoma (A549), and melanoma (G361).

Example 6: Selective interaction of BBP1 with human BAP versus rodent BAP

There are three amino acid substitutions (G5R, F10Y and R13H) in the rodent BAP sequence compared to the human sequence. The rodent peptide demonstrated reduced neurotoxicity and an absence of binding to human brain homogenates (Maggio et al., 1992). It was of interest, therefore, to evaluate the association of rodent BAP with BBP1 in the Y2H system. The sequence of human BAP in pEK162 was changed to encode the rodent peptide by oligonucleotide directed mutagenesis by PCR, described above. The resultant plasmid, pEK240, was identical to the human BAP fusion protein expression plasmid utilized throughout the present invention except for the three codons producing the amino acid substitutions for the rodent peptide sequence. Interactions between BBP1 fusion protein and rodent and human BAP fusion proteins were compared by Y2H bioassay. Strains expressing BBP1 and the rodent BAP failed to produce a growth response (Figure 7). This finding supports the premise that BBP1 may serve as a specific mediator of the neurotoxic effects of BAP, and provides a mechanism to explain the reduced neurotoxicity of the rodent BAP. Importantly, these data also serve to illustrate the high degree of specificity of the BBP1/BAP interaction in the Y2H assays since the substitution of three amino acids was sufficient to completely abrogate the association (Figure 7).

30 Example 7: In vitro binding of radiolabeled beta-amyloid protein to BBP1 protein

Initially, the novel gene product, BBP1, expressed from a fetal brain library as a fusion protein, was shown to interact with beta-amyloid protein (BAP), also expressed as a fusion protein via a yeast 2 hybrid system. To confirm these initial findings, the potential binding of beta-
5 amyloid protein to full-length BBP1 protein was investigated in an in vitro radioligand binding assay. Specifically, radiolabeled human beta-amyloid protein (1-42) was shown to bind with in vitro synthesized myc-tagged BBP1 protein, as evidenced by the ability to co-precipitate beta-amyloid protein with tagged-BBP1 protein. The details of the radioligand binding
10 assay are described below.

Protein A agarose bead + secondary antibody complexes were generated by incubating 2.5µL ImmunoPurePlus immobilized Protein A (Pierce, Rockford, IL) with 10 mg AffiniPure rabbit a-mouse IgG (H + L) (Jackson ImmunoResearch Laboratories, Inc., West Grove, PA) in 50mL
15 cold low salt binding buffer (50mM Tris pH7.6, 150mM NaCl, 2 mM EDTA 1%IGEPAL, and protease inhibitors [5µg/mL leupeptin, 5µg/mL aprotinin, 2µg/mL pepstatin A, 0.25mMPMSF]) with rotation overnight at 4° C. The beads were washed 4X with 1mL binding buffer and were resuspended in 1.25mL binding buffer to give a 50% slurry. In some
20 experiments, a 250mL aliquot of this slurry was incubated in Superblock (Pierce) with rotation overnight at 4° C. The beads were washed 4X with 1mL Superblock and resuspended in 125µL Superblock.

The DNA template for in vitro transcription/translation of the BBP1 protein, including a Kozak consensus sequence and sequences encoding a
25 myc epitope, EQKLISEEDL, directly upstream of the first methionine of BBP1 coding region, was inserted into the BamHI/EcoRI sites of pSP64polyA vector (Promega, Madison, WI). The DNA template was, in part, PCR generated, utilizing the forward primer, 5'
GCAGGATCCCCACCATGGAGCAGAAGCTGATCAGCGAGGAGGACCTGC
30 ATATTTAAAAGGGTCTCCAATGTGA 3' and reverse primer, 5'
TCACGGCCTCCGGAGCAGACGG 3' and PFU polymerase, according to

the manufacturer's conditions (Stratagene, La Jolla, CA). The PCR cycling conditions were an initial denaturing step at 95°C for 3min, 30 cycles of denaturation at 94°C for 30 sec, annealing at 65°C for 30 sec, elongation at 72°C for 1min 30sec, and followed by a final elongation at 72°C for 5 min. The amplicon was digested with BamHI + NotI and ligated to the 3' end of BBP1, housed on a NotI/EcoRI fragment, which had been previously gel purified from the recombinant expression cassette.

Approximately 2.5 μ Ci of disaggregated [Krishnamurthy, K. et al.

10 (1998). Characterization of fibrillogenesis of amyloid peptide. Abstracts of the American Chemical Society, vol. , p. , 215th National Meeting and Exposition, March 29-April 2, 1998, Dallas, Tx.] human [¹²⁵I]-Tyr-Ab₍₁₋₄₂₎ (American Radiolabeled Chemicals, Inc., St. Louis, MO) was incubated with 5-10mL of N-terminal c-myc tagged human BBP1 (1/5-1/10 reaction 15 volume obtained using the TNT SP6 Coupled Reticulocyte Lysate System [Promega, Madison, WI]) with rotation for ~6hrs at 4⁰C in a final volume of 1mL cold low salt binding buffer (see above). Two micrograms of mouse a-myc and 25mL of the Agarose protein A/rabbit a-mouse IgG complex (see above) were added to the reaction tube and incubated at 4⁰ 20 C overnight with rotation. Immune complexes were washed 4X with 1mL binding buffer and resuspended in 25mL 2X Tricine loading dye (Novex, San Diego, CA) containing 5% b-Mercaptoethanol. Samples were boiled for 5 minutes and immediately placed on ice for 15 minutes. The tubes were briefly spun at 2500 x g and the supernatant loaded on a 16% 25 Tricine polyacrylamide gel (Novex, San Diego, CA) which was run at 50mA for ~90 min. The gel was soaked for 15 min in a drying solution composed of 20% acetic acid/10% methanol and dried at 80⁰C for 1 hr under vacuum. The dried gel was subjected overnight to a phosphoimager screen which was scanned for analysis with the Storm 30 phosphoimager (Molecular Dynamics, Sunnyvale, CA).

Initial experiments attempting to co-immunoprecipitate radiolabeled BAP with myc-tagged BBP1 resulted in nonspecific binding of BAP when agarose protein A/secondary antibody complexes were prepared in low salt binding buffer, even in samples lacking BBP1 (see Fig. 8, lane 7). To 5 reduce these non-specific interactions, the agarose protein A/rabbit α-mouse IgG was incubated/washed in blocking reagent prior to binding, as outlined above. This blocking procedure reduced non-specific Ab binding to near zero when all immunoprecipitation components were available except myc-tagged BBP1 (Fig. 8, lane 5). Radiolabeled human BAP₍₁₋₄₂₎ 10 was able to complex with in vitro transcribed/translated myc-tagged human BBP1 after immunoprecipitating myc-tagged BBP1 with anti-myc antibody (Fig. 8, lane 9), as seen by a band consistent in size with Ab (Fig. 8, lane 3). These data are consistent with human BAP binding to myc-tagged human BBP1 in vitro and support the initial observation that 15 BAP interacts with BBP1 in a yeast two-hybrid system.

Example 8: Expression of recombinant BBP1 sensitizes Ntera2 stem cells to β-amyloid peptide.

A cultured cell system was utilized to investigate the effects of 20 BBP1 expression on cellular sensitivity to BAP toxicity. Human Ntera-2 (Nt2) stem cells can be induced to differentiate into neuron-like cells (P. Andrews, Dev. Biol. 103:285-293, 1984). In that state, the cells exhibit a vulnerability to BAP that is similar in degree to that observed in primary neurons. Neurons affected by BAP exhibit characteristics of apoptosis 25 before dying (see C. Cotman and A. Anderson, Mol. Neurobiol. 10:19-45, 1995). An early indicator of apoptosis, namely, condensation of chromatin, was used as an indicator for cellular responses to BAP. The undifferentiated stem cells did not exhibit significant sensitivity under the experimental conditions used in these studies. However, Nt2 stem cells 30 transfected with a BBP1 expression plasmid became markedly sensitive to applied BAP, supporting the premise that BBP1 may act as a mediator of

the toxic effects of b-amyloid peptide. The details of the experiment are below.

BBP cDNAs were modified by polymerase chain reaction (PCR) for expression from the vector pcDNA3.1 (Invitrogen Corp., Carlsbad, CA).

5 BBP1 cDNA was amplified from pBBP1-fl, adding a 5' EcoRI and a 3' Sall site for cloning. The PCR primers were 5' - TGGTGAATTC
GAAAGTGTG GTCTCCAAG ATG G (+ strand) and 5' - CTTCGTCGAC
TTA TGG ATA TAA TTG CGT TTT TC (- strand). The PCR product was
digested with EcoRI + Sall and cloned into pcDNA3.1/EcoRI-Xhol to
10 create pOZ363. Mutation of the arginine codon within the 'DRF' motif of
the BBP1 cDNA was performed using the QuickChange system
(Stratagene Co., La Jolla, CA). Oligonucleotides were synthesized and
purified by Genosys Biotechnologies, Inc. (The Woodlands, TX). The
R138 codon of BBP1 in pOZ363 was changed to a glutamate codon using
15 the oligonucleotide 5' - GG TTG GGA GCA GAT GAA TTT TAC CTT GGA
TAC CC and its exact reverse complement.

Human Ntera2 (Nt2) stem cells were maintained in Dulbecco's
Modified Eagle's medium (high glucose) supplemented with 10% fetal
bovine serum. Retinoic acid was utilized to differentiate cells to a
20 neuronal phenotype as described by P. Andrews (Dev. Biol. 103:285-293,
1984). Expression constructs were introduced into stem cells by
electroporation. The cells were split 1:2 the day before electroporation to
ensure exponential growth for maximal survival and efficiency. On the
day of electroporation the cells were treated with trypsin and washed two
25 times in phosphate buffered saline (PBS). They were resuspended at
 1.3×10^7 cells per 0.3 ml in RPMI 1640 with 10 mM dextrose and 0.1 mM
dithiothriitol. DNA amounts were 7.5 mg subject DNA with 2.5 mg
pEGFP-N1 (CLONTECH Laboratories, Palo Alto, CA) to monitor
transfection. Cells were pre-incubated for 10 mins on ice with DNA,
30 pulsed, and post-incubated for 10 min on ice. A GenePulser instrument
(BioRad Corp., Hercules, CA) was utilized with a cuvette gap of 0.4 cm,

voltage of 0.24 kV, and capacitance of 960 mF. Cells were plated in standard 24-well plates. Approximately 24 hrs after transfection, growth medium was replaced with medium containing the indicated concentration of BAP. After incubation for 44 to 48 hrs, the chromatin-specific dye

5 Hoechst 33342 (Molecular Probes, Inc., Eugene, OR) was added to a concentration of 10 ng/ml. Medium was removed after 10 min and cells were washed with PBS. Cells were then fixed by immersion in PBS containing 4% paraformaldehyde.

Forty-residue β -amyloid peptide was obtained from AnaSpec, Inc.,

10 San Jose, CA. Peptide was dissolved and stored in hexafluoro-isopropanol at 1 mg/ml. Peptide was lyophilized by pervasion with nitrogen, then resuspended in 1.155 ml cell growth medium and divided into 0.13 ml aliquots in a 96-well plate. The plate was shaken at 500 rpm for 4 hrs. Samples were then combined and normalized to a final 15 BAP concentration of 50 mM. The same preparation of aggregated (or aged) BAP utilized in the described experiments was also shown to be toxic to primary hippocampal neurons (data not shown). Forty-two residue β -amyloid peptide was obtained from Bachem Bioscience Inc. It was dissolved directly in cell growth medium and added to experimental 20 samples. This preparation had no discernible effect on differentiated Nt2 neurons.

Cells were visualized on a Zeiss Axiovert fluorescent microscope fitted with dichroic filters as follows. Hoechst dye visualization utilized excitation at 330 microns, emission at 450; EGFP visualization with excitation at 475, emission at 535. A minimum of 60 transfected (EGFP+) cells were scored per sample.

β -amyloid peptide exhibited substantial neurotoxicity in culture only after aging to produce fibrillar aggregates. Peptide freshly dissolved in media showed reduced potency. To investigate potential BBP1 effects on BAP-mediated toxicity, Nt2 stem cells were transfected with pEGFP or with pEGFP plus the BBP1 expression plasmid pOZ363 as described.

These samples were treated with aged (toxic) or fresh (nontoxic) BAP as described above. Nt2 neurons, which exhibited a much greater sensitivity to BAP than the undifferentiated stem cells, were treated at the same time to assess toxicity of the BAP preparations. After treatment for 48 hours, the chromatin of cells was stained with Hoechst dye to reveal nuclear structure and the cells were then fixed. The neurons treated with aged BAP exhibited a marked increase in condensed nuclei compared to those treated with the fresh BAP (Fig. 9), confirming the respective toxicity of the two preparations. In contrast to the observations with Nt2 neurons, the stem cell controls exhibited only a small (not statistically significant) response to the toxic BAP (pEGFP; Fig. 9). However, transfection of the cells with the BBP1 expression plasmid resulted in a significant ($P < 0.001$, Yates modified chi-squared test) increase in the frequency of condensed nuclei (Fig. 9). Importantly, the nontoxic preparation of BAP had no effect in the assay (Fig. 9), establishing that the BBP1-mediated response in these assays is specific for formulations of BAP that are also toxic to Nt2 neurons, and suggesting a correlation between BBP1 and the mechanism of BAP toxicity. In a similar experiment, the toxic BAP was added to transfected Nt2 stem cells at varying concentrations (0, 0.5, 2, or 8 micromolar). Again, cells transfected with pEGFP alone exhibited no significant response to BAP treatment (Fig. 10). In contrast, cells transfected with a BBP1 expression plasmid demonstrated a substantial, significant ($P < 0.005$), and dose-dependent sensitivity to the toxic BAP (Fig. 10).

It was predicted that the BBP1 protein might modulate the activity of heterotrimeric G proteins based on its structural relationship to known G protein-coupled receptors and the demonstration that its predicted cytosolic loop can associate with Ga proteins in a yeast 2-hybrid assay (Example 3, Fig. 3). It has been shown that substitution of the conserved arginine residue in the 'DRF' motif of 7-transmembrane domain G protein-coupled receptors attenuates their activity (E. Burstein, T. Spalding, and

M. Brann, J. Biol. Chem. 273:24322-24327, 1998; W. Rosenthal et al.,

J. Biol. Chem. 268:13030-13033, 1993), and that substitution to

glutamate (E) can completely eliminate agonist-mediated activation of G

protein (P. Jones, C. Curtis, and E. Hulme, Eur. J. Pharmacol. 288:251-

5 257, 1995). The BBP1 arginine-138 codon was mutated to a codon for glutamate and this BBP1 variant was examined for induction of sensitivity to BAP. This single amino acid substitution abrogated the effects of BBP1 (Fig. 10), suggesting that the BAP sensitization affected by BBP1 in this system may be mediated through a G protein pathway.

10 It is clear that the invention may be practiced otherwise than as particularly described in the foregoing description and examples. Numerous modifications and variations of the present invention are possible in light of the above teachings and therefore are within the scope of the appended claims.

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SEQUENCE LISTING

(1) GENERAL INFORMATION:

(i) APPLICANT: Ozenberger, Brad A.
Jacobsen, J. S. Jonathan Bard
Kajkowski, Eileen Steven Walker

(ii) TITLE OF INVENTION: β -Amyloid Peptide-Binding Proteins
and Polynucleotides Encoding the Same

(iii) NUMBER OF SEQUENCES: 2

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(v) COMPUTER READABLE FORM:
(A) MEDIUM TYPE: Floppy disk
(B) COMPUTER: IBM PC compatible
(C) OPERATING SYSTEM: PC-DOS/MS-DOS
(D) SOFTWARE: PatentIn Release #1.0, Version #1.30

(vi) CURRENT APPLICATION DATA:
(A) APPLICATION NUMBER: US
(B) FILING DATE:
(C) CLASSIFICATION:

(viii) ATTORNEY/AGENT INFORMATION:
(A) NAME: Walsh, Andrea C.
(B) REGISTRATION NUMBER: 34,988
(C) REFERENCE/DOCKET NUMBER: 98126

(ix) TELECOMMUNICATION INFORMATION:
(A) TELEPHONE: 973-683-2169
(B) TELEFAX: 973-683-4117

(2) INFORMATION FOR SEQ ID NO:1:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 810 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: mRNA

(ix) FEATURE:

- (A) NAME/KEY: CDS
- (B) LOCATION: 1..807

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

ATG CAT ATT TTA AAA GGG TCT CCC AAT GTG ATT CCA CGG GCT CAC GGG	48
Met His Ile Leu Lys Gly Ser Pro Asn Val Ile Pro Arg Ala His Gly	
1 5 10 15	
CAG AAG AAC ACG CGA AGA GAC GGA ACT GGC CTC TAT CCT ATG CGA GGT	96
Gln Lys Asn Thr Arg Arg Asp Gly Thr Gly Leu Tyr Pro Met Arg Gly	
20 25 30	
CCC TTT AAG AAC CTC GCC CTG TTG CCC TTC TCC CTC CCG CTC CTG GGC	144
Pro Phe Lys Asn Leu Ala Leu Pro Phe Ser Leu Pro Leu Leu Gly	
35 40 45	
GGA GGC GGA AGC GGA AGT GGC GAG AAA GTG TCG GTC TCC AAG ATG GCG	192
Gly Gly Ser Gly Ser Gly Glu Lys Val Ser Val Ser Lys Met Ala	
50 55 60	
GCC GCC TGG CCG TCT GGT CCG TCT GCT CCG GAG GCC GTG ACG GCC AGA	240
Ala Ala Trp Pro Ser Gly Pro Ser Ala Pro Glu Ala Val Thr Ala Arg	
65 70 75 80	
CTC GTT GGT GTC CTG TGG TTC GTC TCA GTC ACT ACA GGA CCC TGG GGG	288
Leu Val Gly Val Leu Trp Phe Val Ser Val Thr Thr Gly Pro Trp Gly	
85 90 95	
GCT GTT GCC ACC TCC GCC GGG GGC GAG GAG TCG CTT AAG TGC GAG GAC	336
Ala Val Ala Thr Ser Ala Gly Gly Glu Ser Leu Lys Cys Glu Asp	
100 105 110	

CTC AAA GTG GGA CAA TAT ATT TGT AAA GAT CCA AAA ATA AAT GAC GCT		384
Leu Lys Val Gly Gln Tyr Ile Cys Lys Asp Pro Lys Ile Asn Asp Ala		
115	120	125
ACG CAA GAA CCA GTT AAC TGT ACA AAC TAC ACA GCT CAT GTT TCC TGT		432
Thr Gln Glu Pro Val Asn Cys Thr Asn Tyr Thr Ala His Val Ser Cys		
130	135	140
TTT CCA GCA CCC AAC ATA ACT TGT AAG GAT TCC AGT GGC AAT GAA ACA		480
Phe Pro Ala Pro Asn Ile Thr Cys Lys Asp Ser Ser Gly Asn Glu Thr		
145	150	155
CAT TTT ACT GGG AAC GAA GTT GGT TTT TTC AAG CCC ATA TCT TGC CGA		528
His Phe Thr Gly Asn Glu Val Gly Phe Phe Lys Pro Ile Ser Cys Arg		
165	170	175
AAT GTA AAT GGC TAT TCC TAC AAA GTG GCA GTC GCA TTG TCT CTT TTT		576
Asn Val Asn Gly Tyr Ser Tyr Lys Val Ala Val Ala Leu Ser Leu Phe		
180	185	190
CTT GGA TGG TTG GGA GCA GAT CGA TTT TAC CTT GGA TAC CCT GCT TTG		624
Leu Gly Trp Leu Gly Ala Asp Arg Phe Tyr Leu Gly Tyr Pro Ala Leu		
195	200	205
GGT TTG TTA AAG TTT TGC ACT GTA GGG TTT TGT GGA ATT GGG AGC CTA		672
Gly Leu Leu Lys Phe Cys Thr Val Gly Phe Cys Gly Ile Gly Ser Leu		
210	215	220
ATT GAT TTC ATT CTT ATT TCA ATG CAG ATT GTT GGA CCT TCA GAT GGA		720
Ile Asp Phe Ile Leu Ile Ser Met Gln Ile Val Gly Pro Ser Asp Gly		
225	230	235
AGT AGT TAC ATT ATA GAT TAC TAT GGA ACC AGA CTT ACA AGA CTG AGT		768
Ser Ser Tyr Ile Ile Asp Tyr Tyr Gly Thr Arg Leu Thr Arg Leu Ser		
245	250	255
ATT ACT AAT GAA ACA TTT AGA AAA ACG CAA TTA TAT CCA TAA		810
Ile Thr Asn Glu Thr Phe Arg Lys Thr Gln Leu Tyr Pro		
260	265	

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 269 amino acids
- (B) TYPE: amino acid
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Met His Ile Leu Lys Gly Ser Pro Asn Val Ile Pro Arg Ala His Gly			
1	5	10	15
Gln Lys Asn Thr Arg Arg Asp Gly Thr Gly Leu Tyr Pro Met Arg Gly			
20	25	30	
Pro Phe Lys Asn Leu Ala Leu Leu Pro Phe Ser Leu Pro Leu Leu Gly			
35	40	45	
Gly Gly Gly Ser Gly Ser Gly Glu Lys Val Ser Val Ser Lys Met Ala			
50	55	60	
Ala Ala Trp Pro Ser Gly Pro Ser Ala Pro Glu Ala Val Thr Ala Arg			
65	70	75	80
Leu Val Gly Val Leu Trp Phe Val Ser Val Thr Thr Gly Pro Trp Gly			
85	90	95	
Ala Val Ala Thr Ser Ala Gly Gly Glu Ser Leu Lys Cys Glu Asp			
100	105	110	
Leu Lys Val Gly Gln Tyr Ile Cys Lys Asp Pro Lys Ile Asn Asp Ala			
115	120	125	
Thr Gln Glu Pro Val Asn Cys Thr Asn Tyr Thr Ala His Val Ser Cys			
130	135	140	
Phe Pro Ala Pro Asn Ile Thr Cys Lys Asp Ser Ser Gly Asn Glu Thr			
145	150	155	160
His Phe Thr Gly Asn Glu Val Gly Phe Phe Lys Pro Ile Ser Cys Arg			
165	170	175	

Asn Val Asn Gly Tyr Ser Tyr Lys Val Ala Val Ala Leu Ser Leu Phe
180 185 190

Leu Gly Trp Leu Gly Ala Asp Arg Phe Tyr Leu Gly Tyr Pro Ala Leu
195 200 205

Gly Leu Leu Lys Phe Cys Thr Val Gly Phe Cys Gly Ile Gly Ser Leu
210 215 220

Ile Asp Phe Ile Leu Ile Ser Met Gln Ile Val Gly Pro Ser Asp Gly
225 230 235 240

Ser Ser Tyr Ile Ile Asp Tyr Tyr Gly Thr Arg Leu Thr Arg Leu Ser
245 250 255

Ile Thr Asn Glu Thr Phe Arg Lys Thr Gln Leu Tyr Pro
260 265

What is claimed is:

1. An isolated polynucleotide selected from the group consisting of:
 - (a) a polynucleotide comprising the nucleotide sequence of SEQ ID NO: 1;
 - (b) a polynucleotide comprising the nucleotide sequence of a β -amyloid peptide-binding protein (BBP) of clone BBP1-fl deposited under accession number ATCC 98617;
 - (c) a polynucleotide encoding a β -amyloid peptide-binding protein (BBP) encoded by the cDNA insert of clone BBP1-fl deposited under accession number ATCC 98617;
 - (d) a polynucleotide comprising the nucleotide sequence of SEQ ID NO: 1 from nucleotide 202 to nucleotide 807;
 - (e) a polynucleotide comprising the nucleotide sequence of a β -amyloid peptide-binding protein (BBP) of clone pEK196 deposited under accession number ATCC 98399;
 - (f) a polynucleotide encoding a β -amyloid peptide-binding protein (BBP) encoded by the cDNA insert of clone pEK196 deposited under accession number ATCC 98399;
 - (g) a polynucleotide encoding a protein comprising the amino acid sequence of SEQ ID NO: 2;
 - (h) a polynucleotide encoding a protein comprising a fragment of the amino acid sequence of SEQ ID NO: 2 having human β -amyloid peptide binding activity, the fragment comprising the amino acid sequence from amino acid 68 to amino acid 269 of SEQ ID NO: 2;
 - (i) a polynucleotide which is an allelic variant of the polynucleotide of (a)-(f) above;
 - (j) a polynucleotide which encodes a species homologue of the protein of (g)-(h) above; and
 - (l) a polynucleotide capable of hybridizing under stringent conditions to any one of the polynucleotides specified in (a)-(h).

2 The polynucleotide of claim 1 wherein said polynucleotide is operably linked to at least one expression control sequence.

3. A host cell transformed with the polynucleotide of claim 2.

4. The host cell of claim 3 wherein said cell is a prokaryotic or eukaryotic cell.

5. A

polynucleotide of claim 2 which process comprises (a) growing a culture of the host cell of claim 3 in a suitable culture medium; and (b) purifying the protein from the culture medium.

10 6. A protein produced according to the process of claim 5.

7. A protein comprising an amino acid sequence selected from the group consisting of:

(a) the amino acid sequence of SEQ ID NO: 2;

(b) the amino acid sequence of SEQ ID NO: 2 from amino acid

15 68 to amino acid 269;

(c) the amino acid sequence encoded by the cDNA insert of clone BBP1-fl deposited under accession number ATCC 98617; and

(d) fragments of the amino acid sequence of SEQ ID NO: 2
using the amino acid sequence from amino acid 185 to amino acid

20 217 of SEQ ID NO: 2.

8. The protein of claim 7, wherein said protein comprises the amino acid sequence of SEQ ID NO: 2.

9. A fusion protein comprising a BBP1 linked to a heterologous protein or peptide sequence.

25 10. The fusion protein of claim 9 in the BBP1 has the amino acid sequence of SEQ ID NO: 2.

11. An oligonucleotide which encodes an antisense sequence complementary to a portion of BBP1 sequence of SEQ ID NO: 1 and which inhibits expression the BBP1 gene.

30 12. A method for determining a polynucleotide encoding a β -amyloid peptide-binding protein (BBP) in a sample comprising the steps of

(a) hybridizing to a sample a probe specific for said polynucleotide under conditions effective for said probe to hybridize specifically to said polynucleotide; and (b) determining the hybridization of said probe to polynucleotides in the sample, wherein said probe comprises a nucleic acid sequence having a region of 20 or more base pairs at least 90% identical to the polynucleotide sequence of SEQ ID NO: 1.

13. A method for determining a polynucleotide encoding a β -amyloid peptide-binding protein (BBP) in a sample comprising the steps of (a) hybridizing to a sample a probe specific for said polynucleotide under conditions effective for said probe to hybridize specifically to said polynucleotide; and (b) determining the hybridization of said probe to polynucleotides in the sample, wherein said probe comprises a nucleic acid sequence having a region of 20 or more base pairs at least 90% identical to the polynucleotide sequence of the cDNA insert of ATCC 98617 or ATCC 98399.

14. An antibody that binds specifically to a polypeptide comprising a region at least 90% identical in sequence to the amino acid sequence of SEQ ID NO: 2.

15. An antibody that binds specifically to a polypeptide comprising a region at least 90% identical in sequence to the amino acid sequence of the β -amyloid peptide binding protein encoded by the cDNA insert of ATCC 98617.

16. A method for detecting in a sample a polypeptide comprising a region at least 90% identical to the amino acid sequence of SEQ ID NO: 2, said method comprising (a) incubating with a sample a reagent that bind specifically to said polypeptide under conditions effective for specific binding; and (b) determining the binding of said reagent to said polypeptide the sample.

17. A method for detecting in a sample a polypeptide comprising a region at least 90% identical in sequence to the amino acid sequence of the β -amyloid peptide binding protein encoded by the cDNA

insert of ATCC 98617, said method comprising (a) incubating with a sample a reagent that bind specifically to said polypeptide under conditions effective for specific binding; and (b) determining the binding of said reagent to said polypeptide the sample.

5 18. A method for diagnosing a disease characterized by aberrant expression of human β -amyloid peptide (BAP), comprising (a) incubating a sample indicative of the aberrant expression of human β -amyloid peptide with a reagent comprising a polypeptide comprising a region at least 90% identical to the amino acid sequence of SEQ ID NO: 10 2 under conditions effective for specific binding of said reagent to said human β -amyloid peptide; and (b) determining the binding of said reagent to said peptide in the sample.

15 19. A method for diagnosing a disease characterized by aberrant expression of human β -amyloid peptide, comprising (a) incubating a sample indicative of the aberrant expression of human β -amyloid peptide with a reagent comprising a polypeptide comprising a region at least 90% identical to the amino acid sequence of the β -amyloid peptide binding protein encoded by the cDNA insert of ATCC 98617 under conditions effective for specific binding of said reagent to said human β -amyloid 20 peptide; and (b) determining the binding of said reagent to said peptide in the sample.

20. A diagnostic process comprising analyzing for the presence of a polynucleotide of claim 1 in a sample derived from a host.

25 21. A method for identifying compounds which regulate the activity of a β -amyloid peptide binding protein comprising (a) incubating a sample comprising human β -amyloid peptide in a test medium containing said test compound and a reagent comprising a polypeptide comprising a region at least 90% identical to the amino acid sequence of SEQ ID NO: 30 2 under conditions effective for specific binding of said reagent to said human β -amyloid peptide; (b) comparing the binding of said reagent to said peptide in the sample in the presence and absence of said test

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compound; and (c) relating the difference between the binding is step (b) to the test compound regulating the activity of the a β -amyloid peptide binding protein.

22. A method for identifying compounds which regulate the

5 activity of a β -amyloid peptide binding protein comprising (a) incubating a sample comprising human β -amyloid peptide in a test medium containing said test compound and a reagent comprising a polypeptide comprising a region at least 90% identical to the amino acid sequence of the β -amyloid peptide binding protein encoded by the cDNA insert of ATCC 98617

10 under conditions effective for specific binding of said reagent to said human β -amyloid peptide; (b) comparing the binding of said reagent to said peptide in the sample in the presence and absence of said test compound; and (c) relating the difference between the binding is step (b) to the test compound regulating the activity of the a β -amyloid peptide

15 binding protein.

23. A method for the treatment of a patient having need to inhibit β -amyloid peptide accumulation in the brain comprising administering to the patient a therapeutically effective amount of the polypeptide of claim 7.

20 24. A transgenic or chimeric animal comprising the polynucleotide of claim 2.

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Abstract of the Invention

**β-AMYLOID PEPTIDE-BINDING PROTEINS
AND POLYNUCLEOTIDES ENCODING THE SAME**

Novel proteins which bind human β -amyloid peptide, polynucleotides which encode these proteins, and methods for producing these proteins are provided. Diagnostic, therapeutic, and screening methods employing the polynucleotides and polypeptides of the present invention are also provided.

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FIGURE 1

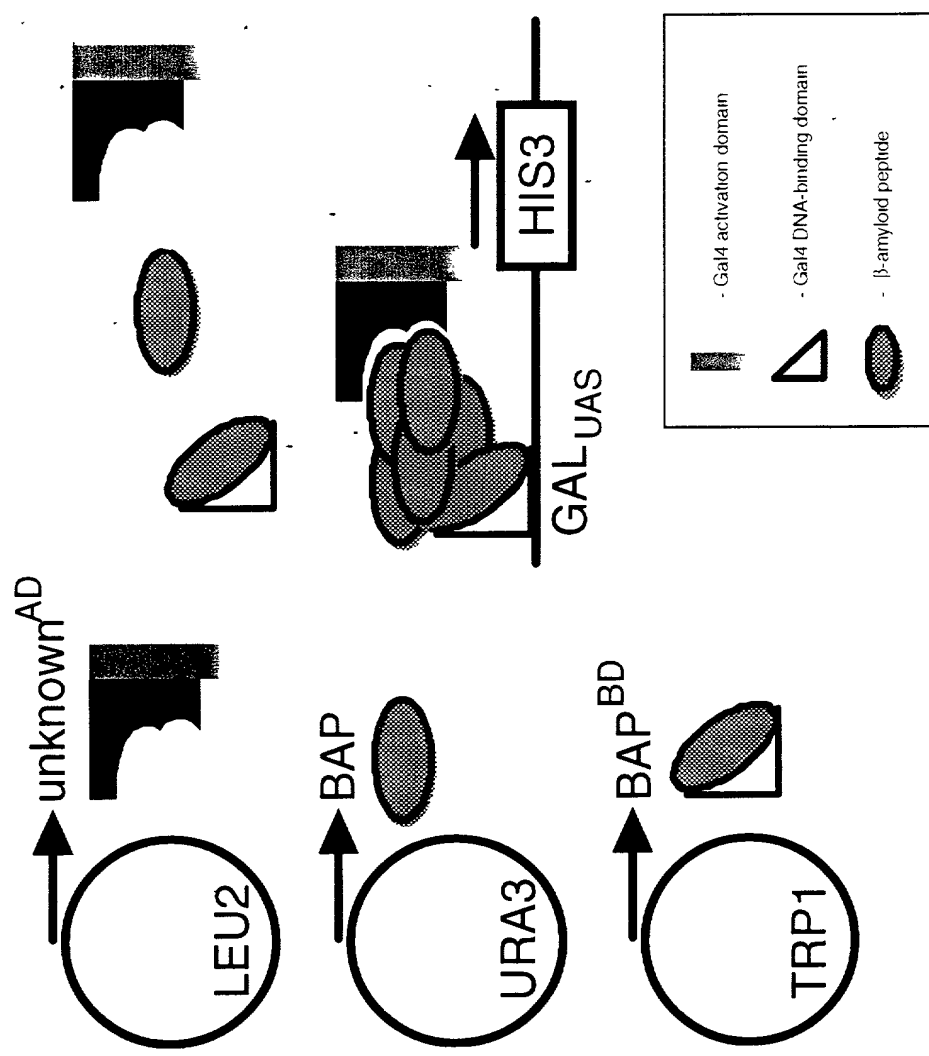


FIGURE 2

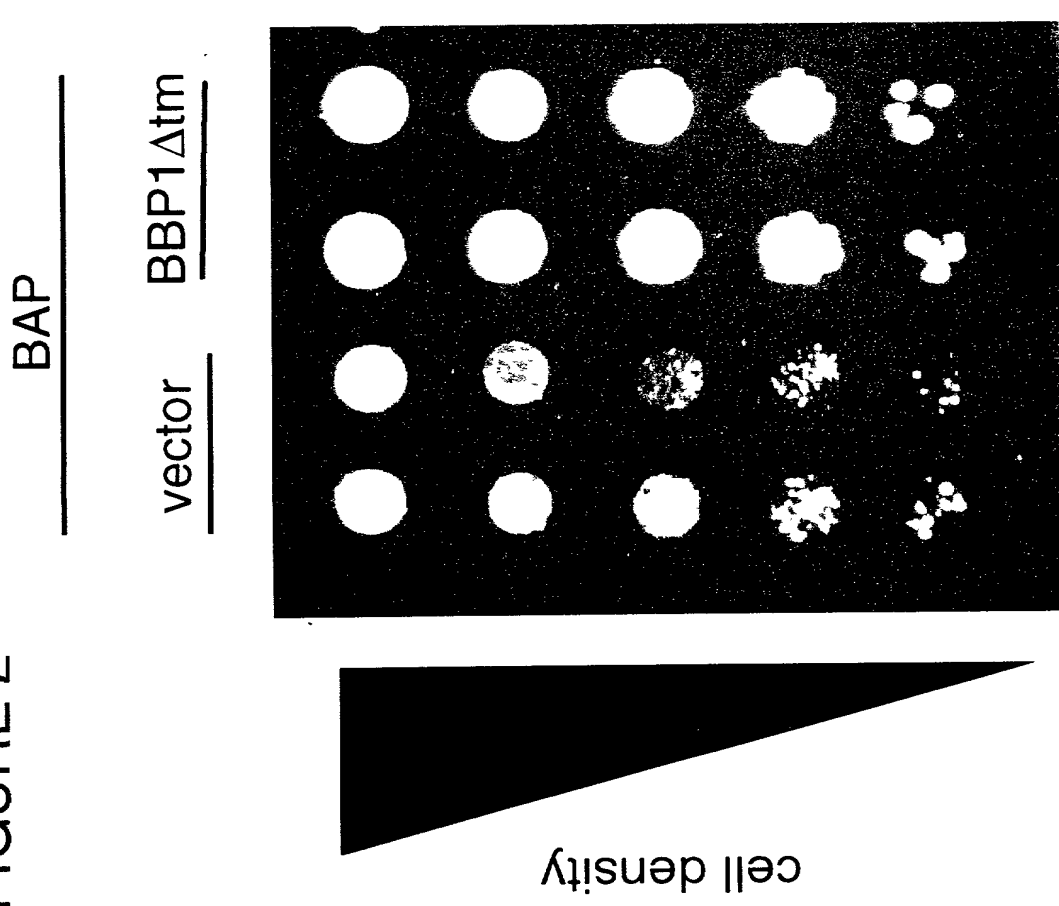


FIGURE 3

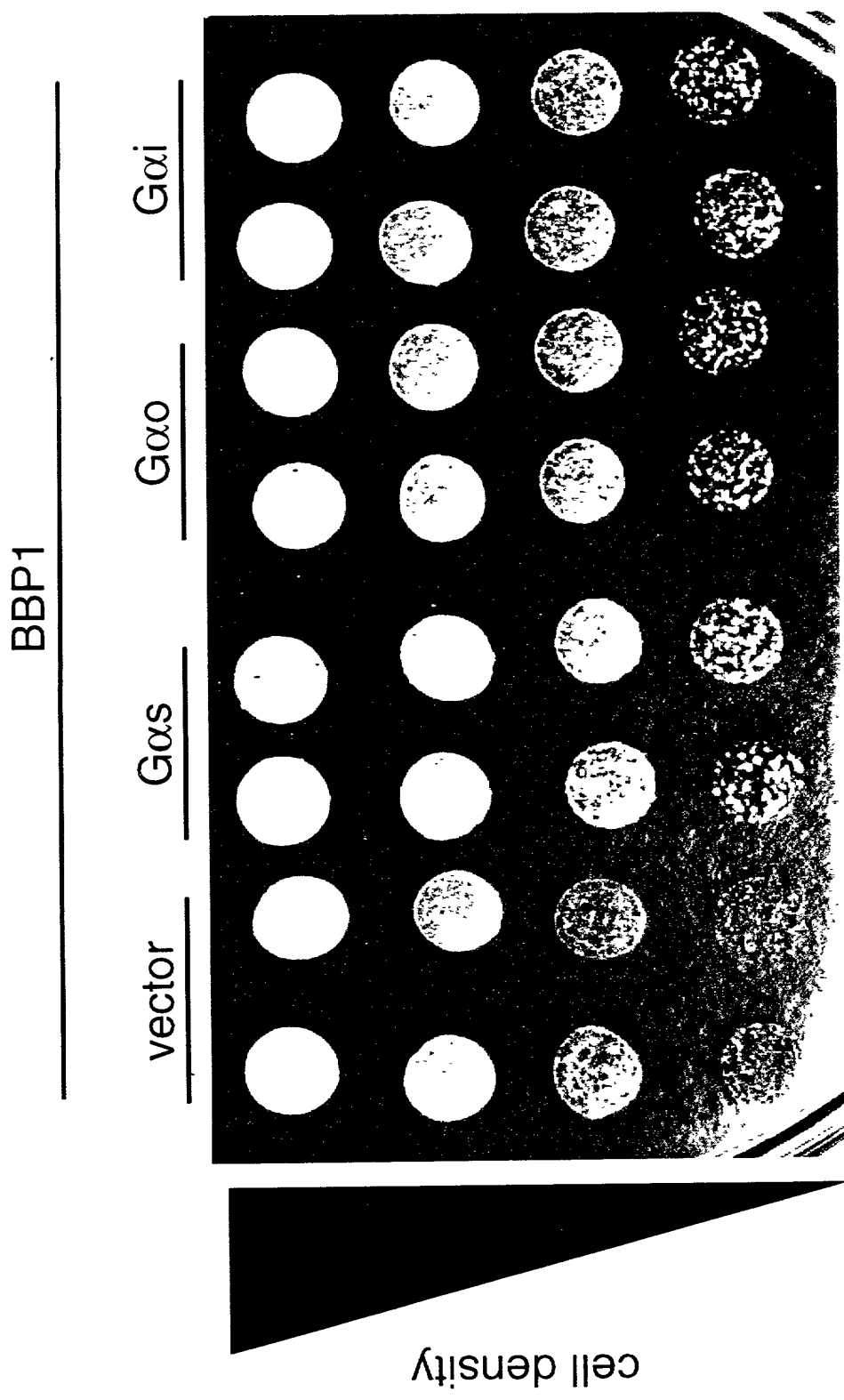


FIGURE 4

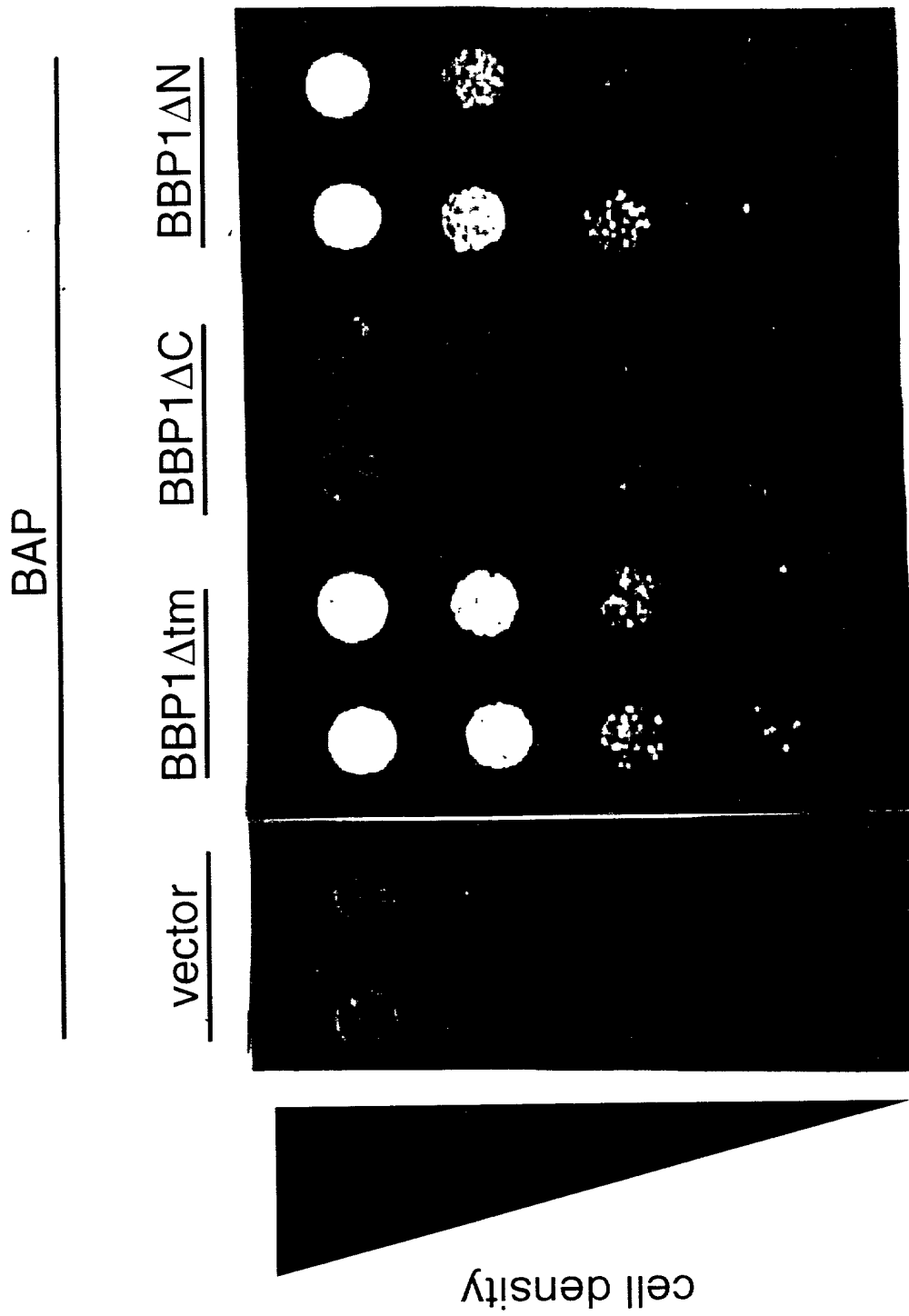


FIGURE 5

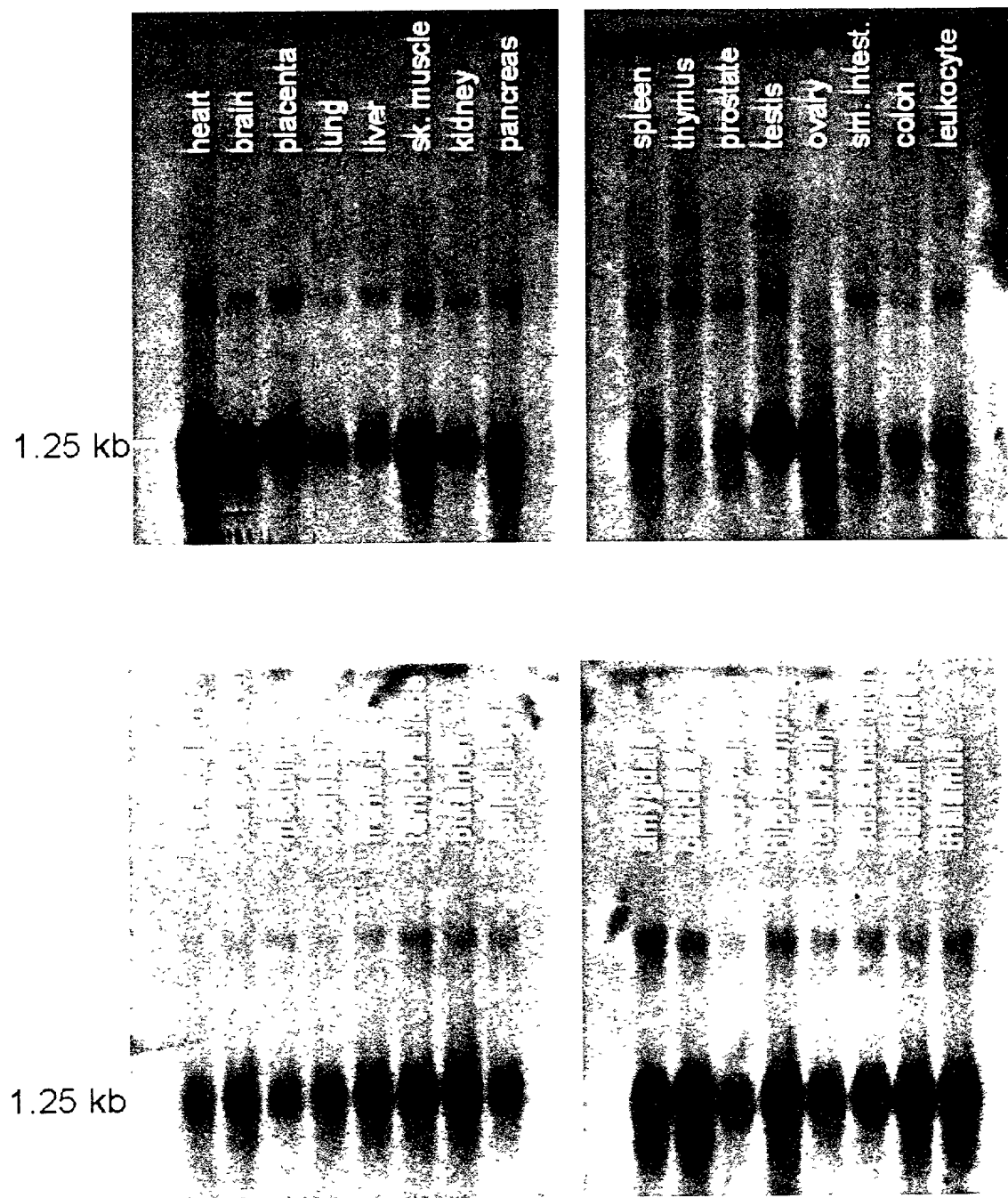


FIGURE 6

A BBP-1



B APP



FIGURE 7

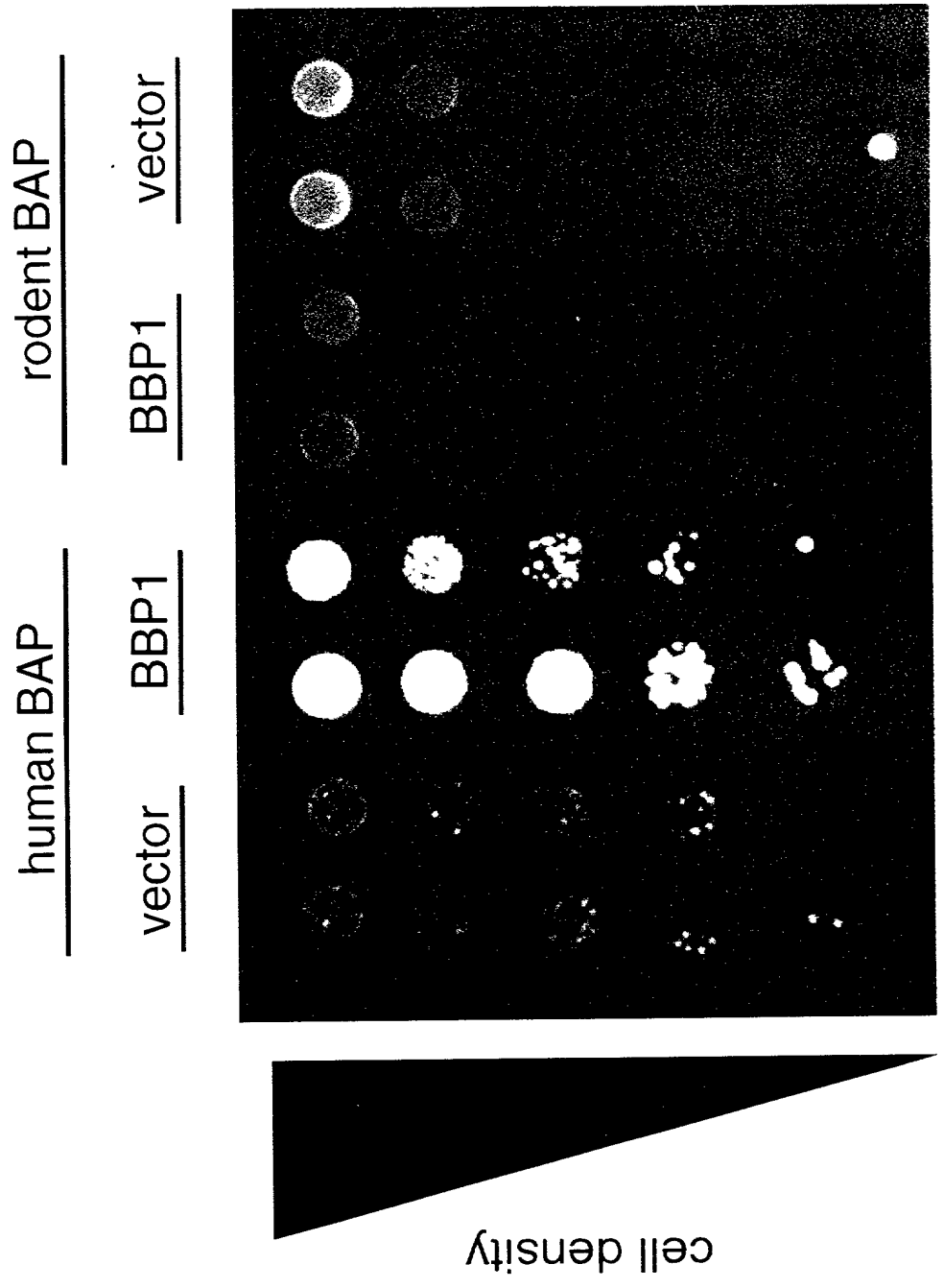


Figure 8

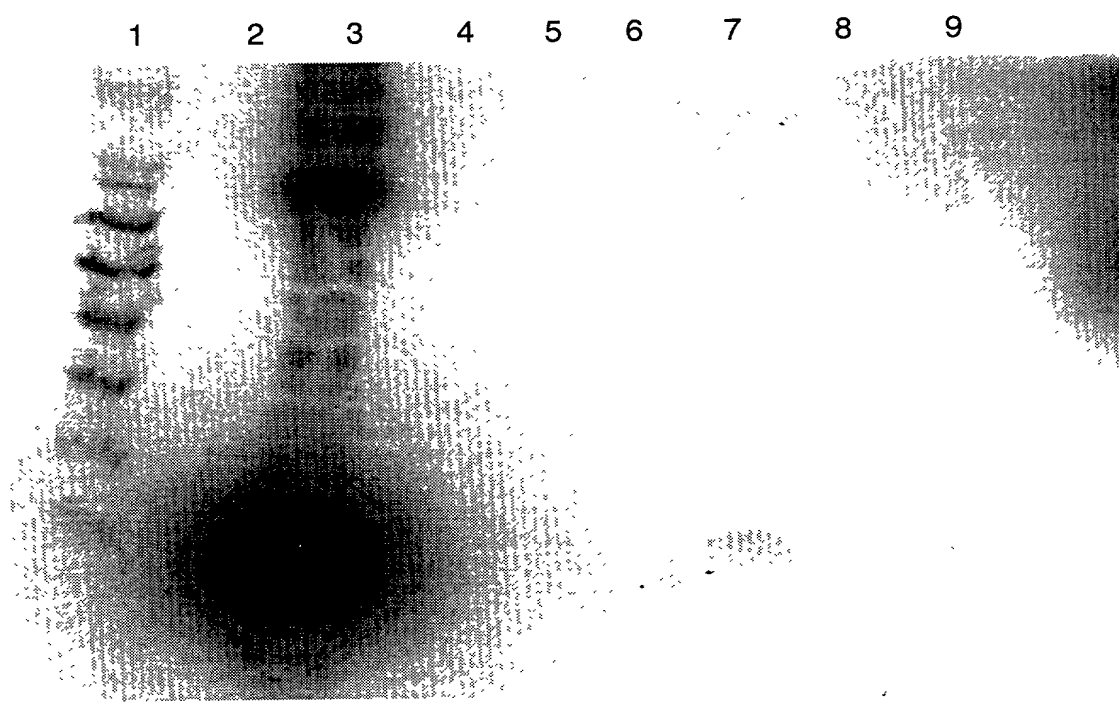


Figure 9

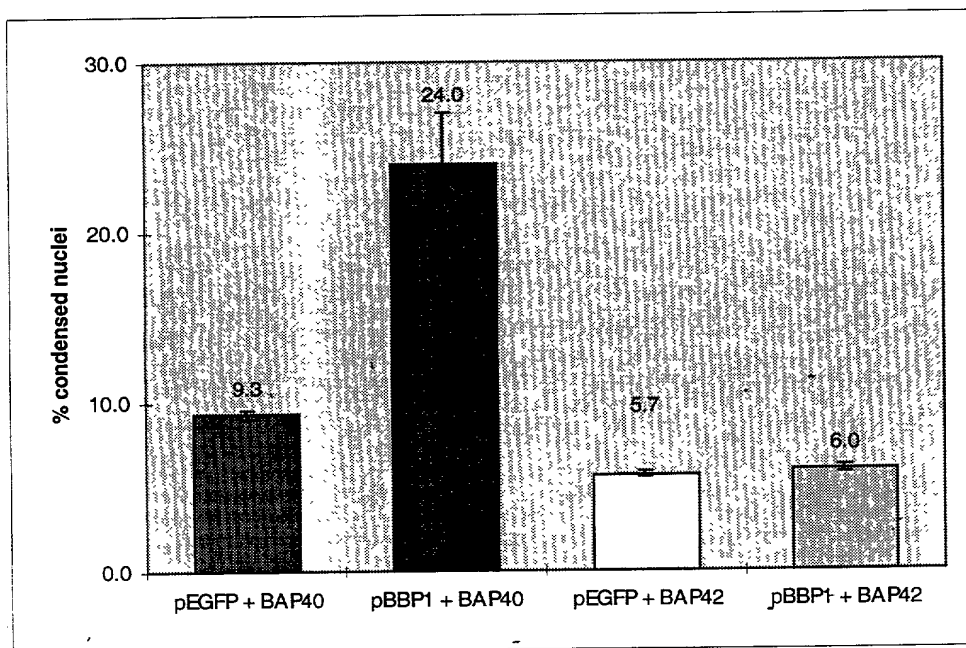
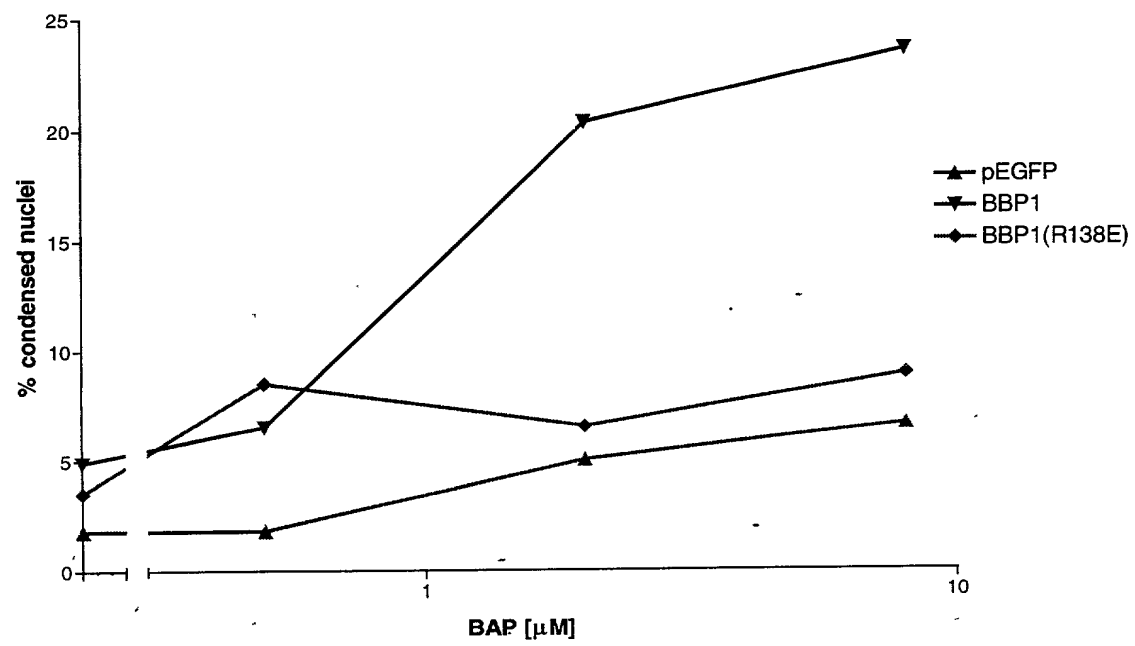


Figure 10



COMBINED DECLARATION AND POWER OF ATTORNEY
(Original, Design, Supplemental, Divisional, Continuation, CIP)

As the below named inventor, I hereby declare that:

TYPE OF DECLARATION

This declaration is of the following type:

- original
- design
- supplemental
- divisional
- continuation
- continuation-in-part (CIP)

INVENTORSHIP IDENTIFICATION

My residence, post office address and citizenship are as stated below next to my name. I believe I am the original, first and sole inventor of the subject matter which is claimed and for which a patent is sought on the invention entitled:

TITLE OF INVENTION

"β-AMYLOID PEPTIDE-BINDING PROTEINS AND POLYNUCLEOTIDES
ENCODING THE SAME"

SPECIFICATION IDENTIFICATION

the specification of which: (complete (a), (b), or (c))

- (a) is attached hereto.
- (b) was filed on as
 - Serial Number
 - Express Mail No. EM165270234US, as Serial Number not yet known
- (c) was described and claimed in PCT International Application No. filed on and as amended under PCT Article 19 on (if any).

ACKNOWLEDGEMENT OF REVIEW OF PAPERS AND DUTY OF CANDOR

I hereby state that I have reviewed and understand the contents of the above identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to disclose information which is material to the examination of this application in accordance with Title 37 CFR 1.56(a).

PRIORITY CLAIM

I hereby claim foreign priority benefits under Title 35, United States Code, Section 119 of any foreign application(s) for patent or inventors certificate or of any PCT International application(s) designating at least one country other than the United States of America listed below and have also identified below any foreign application(s) for patent or inventor's certificate of any PCT International application(s) designating at least one country other than the United States of America filed by me on the same subject matter having a filing date before that of the application(s) of which priority is claimed.

- (d) No such applications have been filed.
- (e) Such applications have been filed as follows.

NOTE: Where item (c) is entered above and the International Application which designated the U.S. claimed priority, check item (e), enter the details below and make the priority claim.

Earliest Foreign Application(s), if any, filed within 12 months (6 months for Design) prior to this U.S. Application

Country	Application No.	Date of Filing (Day, Month, Year)	Priority Claimed 35 USC 119

All Foreign Application(s), if any, Filed More Than 12 Months
(6 Months for Design) Prior to This U.S. Application)

**CLAIM FOR BENEFIT OF PRIOR U.S. PROVISIONAL APPLICATION(S)
(35 U.S.C. § 119(E))**

I hereby claim the benefit under Title 35, United States Code, § 119(e) of any United States provisional application(s) listed below:

PROVISIONAL APPLICATION NUMBER

FILING DATE

60,064,583

4/16/97

POWER OF ATTORNEY

As a named inventor, I hereby appoint the following attorney(s) and/or agent(s) to prosecute this application and transact all business in the Patent and Trademark Office connected therewith.

Andrea C. Walsh	Reg. No. 34,988	Darryl L. Webster	Reg. No. 34,276
Elizabeth M. Barnhard	Reg. No. 31,088	Milagros A. Cepeda	Reg. No. 33,365
Alan M. Gordon	Reg. No. 30,637	Gale F. Matthews	Reg. No. 32,269
		Egon E. Berg	Reg. No. 21,117

Attached as part of this declaration and power of attorney is the authorization of the above-named attorney(s) to accept and follow instructions from my representative(s).

SEND CORRESPONDENCE AND TELEPHONE CALLS TO:

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Patent Law Department- 2B2
One Campus Drive
Parsippany, NJ 07054

Attn: Andrea C. Walsh
Tel. No. (201) 683-2169

CLAIM FOR BENEFIT OF EARLIER U.S./PCT APPLICATION(S) UNDER 35 U.S.C. 120

I hereby claim the benefit under Title 35, United States Code, Section 120 of any United States application(s) designating the United States of America that is/are listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in that/those prior application(s) in the manner provided by the first paragraph of Title 35, United States Code, Section 112, I acknowledge the duty to disclose material information as defined in Title 37, Code of Federal Regulations, Section 1.56(a) which occurred between the filing date of the prior application(s) and the national or PCT International filing date of this application.

**PRIOR U.S. APPLICATIONS OR PCT INTERNATIONAL APPLICATIONS
DESIGNATING THE U.S. FOR BENEFIT UNDER 35 USC 120**

U.S. Applications		Status (Check One)		
U.S. Applications	U.S. Filing Date	Patented	Pending	Abandoned
1 US 60/064,583 - Provisional	4/16/97	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>
2 US 09/060,609	4/15/98	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>
PCT Applications Designating U.S.				
PCT APPLICATION NO.		PCT FILING DATE		U.S. SERIAL NO. ASSIGNED (if any)
3 PCT/US98/07462		4/15/98		
4				

DECLARATION

I hereby declare that all statements herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements are made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application or any patent issued thereon.

